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# VACCINES AGAINST JAPANESE ENCEPHALITIS VIRUS AND WEST NILE VIRUS

#### Field of the Invention

This invention relates to vaccines against Japanese encephalitis virus and West Nile virus.

#### Background of the Invention

The Flavivirus genus of the Flaviviridae family includes approximately 70 viruses, mostly arboviruses, many of which, such as yellow fever (YF), dengue (DEN), Japanese encephalitis (JE), and tick-borne encephalitis (TBE) viruses, are major human pathogens (rev. in Burke and Monath, Fields Virology, 4<sup>th</sup> Ed.:1043-1126, 2001). For example, Japanese encephalitis is the leading cause of viral encephalitis in Asia, where 30,000 to 50,000 new cases are reported each year. As another example, since the first cases were diagnosed in the New York area in 1999, West Nile virus has continued to spread rapidly across North America. The risks of this virus migrating into South America, as well as an epidemic in underdeveloped countries, are extremely high. Effective methods for preventing infection by these viruses are needed, with vaccination being the most cost effective measure.

The Flavivirus particle contains a nucleocapsid composed of viral RNA and capsid protein C. The nucleocapsid is surrounded by an envelope containing the envelope glycoprotein E (50-60 kDa) and a small membrane protein M (7-8 kDa). Translation of the genomic RNA results in a polyprotein precursor that is cleaved by cellular and viral proteases into viral proteins, in the order: C, prM/M, E, NS1, NS2A, NS2B, NS3, NS4A, 2K, NS4B, and NS5, where C through E are the structural components of the virion and NS1 through NS5 are nonstructural proteins required for replication (Lindenbach and Rice, Fields Virology, 4<sup>th</sup> Ed.:991-1041, 2001). The prM protein (~25 kDa) is the intracellular precursor for M. Immature virions containing prM are produced by budding into the lumen of the endoplasmic reticulum (ER) and are transported to the cell surface through the exocytosis pathway. Cleavage of prM

occurs shortly prior to particle release in post-Golgi vesicles. Mature extracellular virus contains predominantly M protein, although a small fraction of uncleaved prM can also be present.

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The E protein is the main functional and antigenic surface component of the virion. The molecular structure of the ectodomain of E, which forms a homodimer on the surface of mature viral particles at neutral pH, has been resolved by cryoelectron microscopy (Rey et al., Nature 375:291-298, 1995) and fitted into the electron density map of viral particles (Kuhn et al., Cell 108:717-725, 2002). During infection, the E protein functions as a class II fusion protein (Modis et al., Nature 427:313-319, 2004). Following virus binding to a cellular receptor and internalization, the acidic pH in the resulting endosomes triggers dissociation of the dimers such that the previously hidden hydrophobic fusion loop of each monomer is exposed outwardly. Concurrently, the loops insert into the cell (endosome) membrane and monomers rearrange into elongated trimers. Further refolding of the trimers brings the cell and viral membranes into close proximity and forces them to fuse, releasing the contents of the viral particle into the cytoplasm. Previous studies showed that some substitutions in the E protein of DEN and JE, which are selected during serial passages in mouse brain and in cultured monkey kidney and mosquito cells, have been localized in particular regions of the 3D structure of the protein, and were reported to be associated with changes in the fusion function of the viruses. The studies showed that the fusion pH threshold for some attenuated vaccines decreased by 0.6 to 1 pH unit by comparison with the corresponding parental virus isolate. Some changes in six residues in the DEN3 protein E (residues 54, 191, 202, 266, 268, and 277) map to the region in domain II. This region is proposed as a focus for the low-pH mediated conformational change required for the surface exposure of the conserved hydrophobic cd fusion loop (Lee et al., Virology 232:281-290, 1997).

There is no evidence that the small (mature) M protein plays a role in the events leading to virus internalization from the endosome or has any other appreciable function, while its intracellular precursor, prM, is known to be important for morphogenesis and transport of progeny viral particles. The prM protein also facilitates proper folding of E (Lorenz et al., J. Virol. 76:5480-5491, 2002) and functions to protect the E protein dimer from premature conformational

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rearrangement during passage of new particles towards the cell surface through acidic secretory compartments (Guirakhoo et al., J. Gen. Virol. 72:1323-1329, 1991; Guirakhoo et al., Virology 191:921-931, 1992).

ChimeriVax<sup>TM</sup> technology has been used to create live, attenuated vaccine candidates against medically important Flaviviruses. It employs the YF 17D vaccine virus as a vector in which the prM-E genes are replaced with the prM-E genes from a heterologous Flavivirus, such as JE, dengue, West Nile, or St. Louis encephalitis viruses (Monath et al., Vaccine 17:1869-1882, 1999; Monath et al., Curr. Drug Targets - Inf. Disorders 1:37-50, 2001; Monath et al., Vaccine 20:1004-1018, 2002; Guirakhoo et al., Virology 257:363-372, 1999; Guirakhoo et al., J. Virol. 75:7290-7304, 2001; Guirakhoo et al., Virology 298:146-159, 2002; Pugachev et al., Int. J. Parasitol. 33:567-582, 2003; Guirakhoo et al., J. Virol. 78:4761-4775, 2004). Previously, the ChimeriVax<sup>TM</sup>-JE vaccine virus, containing the prM-E genes from the SA14-14-2 virus (live attenuated JE vaccine used in China), was propagated to high titers in Vero cells cultured in media supplemented with fetal bovine serum (FBS) (Monath et al., Biologicals 33:131-144, 2005). It was successfully tested in preclinical and Phase I and II clinical trials (Monath et al., Vaccine 20:1004-1018, 2002; Monath et al., J. Infect. Dis. 188:1213-1230, 2003). Similarly, successful Phase I clinical trials have been conducted with a ChimeriVax<sup>TM</sup>-WN vaccine candidate, which contains the prM-E sequence from a West Nile virus (NY99 strain), with three specific amino acid changes incorporated into the E protein to increase attenuation (Arroyo et al., J. Virol. 78:12497-12507, 2004).

#### Summary of the Invention

The invention provides recombinant Flaviviruses that include one or more membrane (M) protein mutations (e.g., substitutions, deletions, or insertions), such as mutations that attenuate the Flavivirus (e.g., mutations that decrease the viscerotropism/viremia of the Flavivirus), increase genetic stability of the Flavivirus during propagation in cell culture (e.g., manufacturing in serum free cultures), and/or increase vaccine virus yields. The Flaviviruses of the invention can be chimeric Flaviviruses, such as Flaviviruses that include capsid and non-structural proteins of a first Flavivirus (e.g., a yellow fever virus, such as YF 17D) and membrane and/or envelope proteins of a second Flavivirus (e.g., Japanese encephalitis virus, West Nile

virus, a dengue virus (dengue-1, dengue-2, dengue-3, or dengue-4 virus), St. Louis encephalitis virus, Murray Valley encephalitis virus, tick-borne encephalitis virus, as well as any other Flavivirus that is a human/animal pathogen from the YF, JE, DEN, and TBE serocomplexes).

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In the Flaviviruses of the invention, the mutation (e.g., substitution) can be in the transmembrane or ectodomain of membrane protein M. For example, the mutation can be in the region of amino acids 40-75 of the predicted membrane helix of the membrane protein M of the Flavivirus. As an example, the mutation can be a substitution of amino acid 60 of the membrane protein of a Flavivirus such as Japanese encephalitis virus (e.g., arginine to cysteine in the Japanese encephalitis virus M protein), or in a corresponding amino acid of another Flavivirus. Determination of which amino acid in a given Flavivirus "corresponds" to that of another Flavivirus can be carried out by standard amino acid sequence alignment, as is well known to those of skill in this art. As another example, the mutation can be a substitution of amino acid 66 of the membrane protein of a Flavivirus such as West Nile virus (e.g., a substitution of leucine with proline in the M protein of West Nile virus), or in a corresponding amino acid of another Flavivirus. In other examples, the mutation is at another membrane anchor amino acid, e.g., one or more amino acids selected from the group flanking the M66 residue, including positions 60, 61, 62, 63, 64, 65, and 66 of Japanese encephalitis virus or West Nile virus (or corresponding amino acids in other Flaviviruses) or other amino acid residues of the transmembrane domain.

We also provide for the first time evidence that the ectodomain of the M protein is of important functional significance, because a glutamine to proline change at the M5 residue increased the pH threshold of infection. Therefore, it can now be expected that Flavivirus attenuation can be achieved through amino acid changes or introduction of various deletions or insertions in the amino-terminal ectodomain, or surface part of the M protein, not only its C-terminal hydrophobic anchor. Thus, in other examples, the viruses of the invention include one or more mutations in the M protein ectodomain (residues 1-40) as described herein. This result is quite unexpected, given the lack of any known function of the mature M protein of Flaviviruses.

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In addition to the membrane protein mutations noted above, in the case of chimeric Flaviviruses that include membrane and envelope proteins of a West Nile virus, the viruses of the invention can include one or more envelope protein mutations in amino acids selected from the group consisting of amino acids 107, 138, 176, 177, 224, 264, 280, 316, and 440. In other Flaviviruses, the mutations can be present in amino acids that correspond to these amino acids. As a specific example, the Flavivirus can include a mutation corresponding to mutation(s) in West Nile M protein amino acid 66 and E protein mutations at amino acids corresponding to West Nile virus amino acids 107, 316, and 440. In addition to the mutations described above, the Flaviviruses of the invention can also include one or more mutations in the hydrophobic pocket of the hinge region of the envelope protein, as described elsewhere herein. Further mutations that can be included in the viruses of the invention are mutations in the 3'UTR, the capsid protein, or other envelope protein regions, as described further below.

The invention also provides vaccine compositions that include the Flaviviruses described above and elsewhere herein and pharmaceutically acceptable carriers or diluents, as well as methods of inducing immune responses to Flaviviruses in patients by administration of such vaccine compositions. The patients treated according to such methods include those that do not have, but are at risk of developing, infection by the Flavivirus, as well as patients that are infected by the Flavivirus. Further, the invention includes the use of the Flaviviruses described herein in the prophylactic and therapeutic methods described herein, as well as in the manufacture of medicaments for these purposes.

The invention further provides methods of producing vaccines that include a Flavivirus as described herein, which involve introducing into the membrane protein of the Flavivirus a mutation that results in decreased viscerotropism/viremia, and/or increased genetic stability/yields. Further, the invention provides nucleic acid molecules (RNA or DNA) corresponding to the genomes of the Flaviviruses described herein (or the complements thereof), and methods of using such nucleic acid molecules to make the viruses of the invention.

The Flaviviruses of the invention are advantageous because, in having decreased virulence (shown, e.g., by decreased viscerotropism/viremia), they provide an additional level of safety, as compared to their non-mutated counterparts, when

administered to patients. An additional advantage is that some mutations, such as the M-60 mutation in ChimeriVax<sup>TM</sup>-JE, preclude accumulation of undesirable mutations during vaccine manufacture that otherwise could compromise safety, and increase manufacturing yields. Additional advantages of these viruses are provided by the fact that they can include sequences of yellow fever virus strain YF17D (e.g., sequences encoding capsid and non-structural proteins), which (i) has had its safety established for >60 years, during which over 350 million doses have been administered to humans, (ii) induces a long duration of immunity after a single dose, and (iii) induces immunity rapidly, within a few days of inoculation. In addition, the vaccine viruses of the invention cause an active infection in the treated patients. As the cytokine milieu and innate immune response of immunized individuals are similar to those in natural infection, the antigenic mass expands in the host, properly folded conformational epitopes are processed efficiently, the adaptive immune response is robust, and memory is established.

The beneficial aspects of mutations in the M protein on vaccine safety and manufacture in cell culture are novel and unexpected, given the lack of any known function of the mature M protein of Flaviviruses.

Other features and advantages of the invention will be apparent from the following detailed description, the drawings, and the claims.

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#### Brief Description of the Drawings

Fig. 1A is a schematic representation of the 3' untranslated region of yellow fever virus, which shows domains within this region (repeat sequences (RS), conserved sequences CS2, CS1, and the 3'-extreme stem-loop structure), as well as examples of mutations that can be included in the viruses of the invention (e.g., deletions dA, dB, dC, dD, d7, d14, CS2 d5, and CS2 d16).

Fig. 1B is a schematic representation of the sequence and published secondary structure prediction of the 3' untranslated region of yellow fever 17D virus, from the middle of the 3<sup>rd</sup> RS element to the end of the UTR (Proutski et al., J. Gen. Virol. 78:1543-1549, 1999).

Fig. 1C is an illustration of the optimal YF 17D 3'UTR secondary structure prediction produced using the Zuker RNA folding algorithm.

Fig. 1D is an illustration of the effects of 3'UTR deletions (shown for the dC deletion; Zuker method) on the optimal YF 17D structure (compare with Fig. 1C).

Fig. 2A is a schematic representation of the sequence of the capsid protein of tick-borne encephalitis virus, as well as deletions in this protein reported by Kofler et al., J. Virol. 76:3534-3543, 2002.

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Fig. 2B is a schematic representation of the sequence of the capsid protein of YF 17D virus. Regions predicted by computer analysis to have α-helical secondary structure (α-helices I-IV), as well as hydrophobic regions (solid bars) and deletions introduced in this protein in certain ChimeriVax<sup>TM</sup>-WN viruses (e.g., deletions C1 and C2; boxed) are indicated.

Fig. 3 is a graph showing growth of the indicated viruses (WN01, WN02 P5, Large Plaque, Small Plaque, and YF/17D) in HepG2 cells.

Fig. 4 is a graph showing growth of the indicated viruses (WN01, WN02 P5, Large Plaque, Small Plaque, and YF/17D) in THLE-3 cells.

Fig. 5 is a graph showing the viremia in hamsters induced by the indicated viruses (WN02 P5; mixed plaque), Small Plaque (PMS, P10), and Large Plaque (PMS, P10)).

Fig. 6 is a schematic representation of the passage of SF ChimeriVax<sup>TM</sup>-JE virus samples (g.s., experimental passages to study genetic stability).

Fig. 7 is a graph showing growth curves of SF ChimeriVax<sup>TM</sup>-JE viruses of the invention (uncloned P2, P3 MS (E-107), P4 PS (E-107), P5 g.s. (M-60), and P5 VB (E-107)) at the indicated times post-infection, which shows higher growth rates in SF culture of virus samples containing the M-60 [arginine (R) → cysteine (C) and E-107 phenylalanine (F)→leucine (L)] mutants as compared to nonmutant virus (P2).

Fig. 8A is a graph showing infectivities of the M-5 ChimeriVax<sup>TM</sup>-JE mutant (Clone E) compared to P5 uncloned vaccine bulk and Clone I (E-107 mutant), non-mutant (Clone A), and M-60 mutant (Clone C) after treatment with a range of acidic pH. Of significance is the appearance of the slopes and at which pH the viruses lost infectivity, but not initial titers in diluted samples (e.g., at pH 6.8).

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Fig. 8B is a Survival Plot of ChimeriVax<sup>TM</sup>-JE vaccine (1.9 log<sub>10</sub> PFU/dose, as determined by back titration of inocula) in comparison to ChimeriVax<sup>TM</sup>-JE M5 mutant (1.4 log<sub>10</sub> PFU/dose, as determined by back titration of inocula) in 3-4 day old suckling mice inoculated by the intracerebral route.

Fig. 8C is a Survival Plot of ChimeriVax<sup>TM</sup>-JE M5 mutant virus (1.4 log<sub>10</sub> PFU/dose as determined by back titration of inocula) in comparison to YF-VAX<sup>®</sup> (0.9 log<sub>10</sub> PFU/dose as determined by back titration of inocula) in 3-4 day old suckling mice inoculated by the intracerebral route.

Fig. 8D shows the results of an Indirect Fusion assay, which provides a comparison of P7 and P10 of ChimeriVax<sup>TM</sup>-DEN1-4 viruses. The virus output for each experiment was determined by standard plaque assay. A, ChimeriVax<sup>TM</sup>-DEN1 PMS P7 (triangles) and P10 (diamonds); B, ChimeriVax<sup>TM</sup>-DEN2 PMS P7 (triangles) and P10 (diamonds); C, ChimeriVax<sup>TM</sup>-DEN3 PMS P7 (triangles) and P10 (diamonds); D, ChimeriVax<sup>TM</sup>-DEN4 PMS P7 (triangles) and P10 (diamonds).

Fig. 8E shows the results of an Indirect Fusion assay with the ChimeriVax<sup>TM</sup>-DEN3, comparing the PMS (P7) vaccine with the Vaccine lot (P10) and the P15 virus. The virus output for each experiment was determined by standard plaque assay. ChimeriVax<sup>TM</sup>-DEN3 PMS P7 (triangles), P10 (diamonds), and P15 (squares).

Fig. 8F shows the structure of a DEN1 E-protein dimer (amino acids 1 to 394)

of ChimeriVax<sup>TM</sup>-DEN1 virus (Guirakhoo et al., J. Virol. 78:9998-10008, 2004). (A)

The position of the positively charged lysine (K) at residue 204 of the P7 (PMS, 204K) virus is shown by CPK (displays spheres sized to van der Waal radii)

representation. Three structural domains are shown in black (domain I), light grey (domain II), and dark grey (domain III). (B) Close up of marked area in panel A. (C)

The same area as in panel B from the E protein model of the mutant DEN1 virus (P10, 204R shown in black). Selected amino acids in panel B and C are shown in stick representation. Medium grey, carbon; dark grey, nitrogen; black, oxygen; light grey, sulfur.

Fig. 9A is a graph showing the penetration efficiency of ChimeriVax<sup>TM</sup>-JE viruses M60 mutant (Clone C), E107 mutant (Clone I), and non-mutant (Clone A) at the indicated times. These results indicate that the M60 mutation facilitates penetration in SF Vero cells apparent at the 5 and 10 minute time points. SF Vero cells were infected with appropriately diluted viruses (Clones A, C, and I in serum

free medium) for 5, 10, 20, or 60 minutes, and then were treated for 3 minutes with a solution of 0.1 M glycine, 0.1 M NaCl, pH 3.0, to inactivate extracellular virus. Wells were washed twice with PBS, and then monolayers were overlaid with methylcellulose followed by staining plaques on day 5 with crystal violet. Efficiency of penetration is shown as percentages of observed plaque numbers after glycine treatment as compared to control infected wells that were treated with PBS instead of glycine.

Fig. 9B is a schematic representation of the locations of the E-107, M-5, and M-60 amino acid residues in the envelope proteins E and M, illustrating the hypothetical effect of the M-5 residue on fusion. The dashed stretch at the tip of domain II of the E protein containing the E-107 residue represents the fusion peptide (c-d loop), which inserts into cell membrane (Rey et al., Nature 375:291-298, 1995). The M-5 residue is in the N terminal part of the ectodomain of the M protein. The E protein monomers rearrange into trimeric complexes, which fold to force the cell and virus membranes to fuse (Modis et al., Nature 427(6972):313-319, 2004). The M protein may be a functional component of the complexes, e.g., facilitating fusion of the viral membrane with the cell membrane via its interaction with the E protein. The M-60 residue is between the two C-terminal transmembrane stretches of M and may participate in the interaction of the cell and viral membranes during fusion.

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#### **Detailed Description**

The invention provides vaccines and methods for use in preventing and treating Flavivirus (e.g., Japanese encephalitis (JE) or West Nile (WN) virus) infection. The methods of the invention generally involve vaccination of subjects with a live, attenuated chimeric Flavivirus that consists of a first Flavivirus (e.g., yellow fever virus) in which one or more structural proteins (e.g., membrane and/or envelope proteins) have been replaced with those of a second Flavivirus (e.g., Japanese encephalitis (JE) and/or West Nile (WN) virus; also see below). The membrane proteins of the chimeras of the invention include one or more mutations, as is described further below. Also as is described below, structural proteins such as membrane and/or envelope proteins of other Flaviviruses can be used in place of those of Japanese encephalitis virus or West Nile virus in the chimeric viruses of the present invention. Further, the membrane protein mutations of the invention can also

be used in intact, non-chimeric Flaviviruses (e.g., any of those listed herein), not including any replacements of structural proteins, and optionally with one or more additional mutations, such as those described herein.

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A specific example of a chimeric virus that can be included in the vaccines of the invention is the human yellow fever vaccine strain, YF 17D (e.g., YF17D-204 (YF-VAX®, Sanofi-Pasteur, Swiftwater, PA, USA; Stamaril®, Sanofi-Pasteur, Marcy-L'Etoile, France; ARILVAX™, Chiron, Speke, Liverpool, UK; FLAVIMUN®, Berna Biotech, Bern, Switzerland); YF17D-204 France (X15067, X15062); YF17D-204, 234 US (Rice et al., Science 229:726-733, 1985)), in which the membrane and envelope proteins have been replaced with the membrane and envelope proteins (including an M protein mutation, such as a substitution in M60, as described herein) of Japanese encephalitis virus. In another example, the YF 17D membrane and envelope proteins are replaced with those of a West Nile virus (including an M protein mutation, such as a substitution in M66, as described herein).

In other examples, another Flavivirus, such as a dengue virus (serotype 1, 2, 3, or 4), St. Louis encephalitis virus, Murray Valley encephalitis virus, yellow fever virus, including YF 17D strains, or any other Flavivirus, can provide the membrane and/or envelope proteins in such a chimeric virus. Additional Flaviviruses that can be attenuated according to the invention, whether as intact, non-chimeric viruses or as the source of membrane and/or envelope proteins in chimeras, include other mosquito-borne Flaviviruses, such as Kunjin, Rocio encephalitis, and Ilheus viruses; tick-borne Flaviviruses, such as Central European encephalitis, Siberian encephalitis, Russian Spring-Summer encephalitis, Kyasanur Forest Disease, Omsk Hemorrhagic fever, Louping ill, Powassan, Negishi, Absettarov, Hansalova, Apoi, and Hypr viruses; as well as viruses from the Hepacivirus genus (e.g., Hepatitis C virus). Other yellow fever virus strains, e.g., YF17DD (GenBank Accession No. U 17066), YF17D-213 (GenBank Accession No. U17067; dos Santos et al., Virus Res. 35:35-41, 1995), and yellow fever virus 17DD strains described by Galler et al., Vaccines 16(9/10):1024-1028, 1998, can also be used as the backbone viruses into which heterologous structural proteins can be inserted according to the invention.

The viruses listed above each have some propensity to infect visceral organs.

The viscerotropism of these viruses may cause dysfunction of vital visceral organs, such as observed in YF vaccine-associated adverse disease events, albeit very

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infrequently. The replication of virus in these organs can also cause viremia and thus contribute to invasion of the central nervous system. Decreasing the viscerotropism of these viruses by mutagenesis according to the present invention can thus reduce the abilities of the viruses to cause adverse viscerotropic disease and/or to invade the brain and cause encephalitis.

The mutations of the invention result in beneficial effects to the viruses, which can include, for example, increased attenuation, stability, and/or replication. The mutations are present in the membrane protein, e.g., in the transmembrane region or in the ectodomain of the membrane protein. For example, the mutations can be in amino acid 60 or 66 of the membrane protein and/or in other amino acids within the predicted transmembrane domain (e.g., in any one or more of amino acids 40-75), or in the N-terminal ectodomain of the M protein (e.g., M-5). As a specific example, membrane protein amino acid 60 (arginine in wild type Japanese Encephalitis virus) can be replaced with another amino acid, such as cysteine. A substitution from arginine to cysteine at position M-60 in the ChimeriVax<sup>TM</sup>-JE virus significantly reduced the viremia (viscerotropism) of the virus for humans in clinical trials in which variants of the vaccine with and without the M-60 mutation were tested (Tables 11A and 11B). In addition to cysteine, other amino acids, such as serine, threonine, glycine, methionine, etc., can substitute the wild type amino acid at position 60 of the membrane protein. In another example, membrane protein amino acid 66 (leucine in wild type West Nile virus) can be replaced with another amino acid, such as proline. In addition to proline, other hydrophobic amino acids, such as isoleucine, methionine, or valine, or small amino acids, such as alanine or glycine, can substitute the wild type amino acid at position 66 of the membrane protein. These mutations can also be present in corresponding amino acids of other Flaviviruses, as described herein.

As other examples of substitutions that can be made in membrane protein sequences, amino acids at positions 61, 62, 63, and/or 64 can be substituted, alone or in combination with each other, a mutation at position 60, a mutation at position 66, and/or another mutation(s). Examples of substitutions at these positions in the West Nile virus membrane protein sequence include: valine to alanine at position 61, valine to glutamic acid or methionine at position 62, phenylalanine to serine at position 63, and valine to isoleucine at position 64. These mutations can also be present in corresponding amino acids of other Flaviviruses, as described herein.

Examples of substitutions at these or surrounding positions in the JE virus membrane protein sequence include any of the remaining 20 amino acids with the expectation that a desired effect on viscerotropism and/or vaccine virus replication/stability in cell culture during manufacturing will be achieved. Other examples in chimeric or non-chimeric Flaviviruses include any amino acid substitutions, alone or in combinations, in the N-terminal ectodomain of the M protein composed of residues 1 - ~40 of the protein, as well as deletion(s) of various sizes (e.g., 1, 2, 3, 4, 5, etc., amino acids long) introduced into the ectodomain and/or the transmembrane domain of the M protein.

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In addition to one or more of the membrane protein mutations noted above, the viruses of the invention can also include one or more additional mutations. For example, in the case of West Nile virus, such an additional mutation(s) can be in the region of position 107 (e.g., L to F), 316 (e.g., A to V), or 440 (e.g., K to R) (or a combination thereof) of the West Nile virus envelope protein. The mutations can thus be, for example, in one or more of amino acids 102-112, 138 (e.g., E to K), 176 (e.g., Y to V), 177 (e.g., T to A), 244 (e.g., E to G), 264 (e.g., Q to H), 280 (e.g., K to M), 311-321, and/or 435-445 of the West Nile envelope protein. As a specific example, using the sequence of West Nile virus strain NY99-flamingo 382-99 (GenBank Accession Number AF196835) as a reference, the lysine at position 107 can be replaced with phenylalanine, the alanine at position 316 can be replaced with valine, and/or the lysine at position 440 can be replaced with arginine. Examples of additional combinations of amino acids that can be mutated include are as follows: 176, 177, and 280; 176, 177, 244, 264, and 280; and 138, 176, 177, and 280. Further, these mutations can also be present in corresponding amino acids of other Flaviviruses, as described herein.

The ChimeriVax<sup>TM</sup>-JE vaccine already includes all of the above-noted SA14-14-2 specific mutations as it contains the SA14-14-2-specific JE envelope. Additional amino acid changes in the E protein can also be selected and introduced based on the knowledge of the structure/function of the E protein for additional attenuation (e.g., as described below). These mutations can also be present in corresponding amino acids of other Flaviviruses, as described herein.

In addition to the amino acids noted above, the substitutions can be made with other amino acids, such as amino acids that would result in conservative changes from those noted above. Conservative substitutions typically include substitutions within the following groups: glycine, alanine, valine, isoleucine, and leucine; aspartic acid, glutamic acid, asparagine, and glutamine; serine and threonine; lysine and arginine; and phenylalanine and tyrosine.

The viruses of the invention (e.g., Japanese encephalitis and West Nile viruses, and chimeric Flaviviruses including membrane and envelope proteins from these or other flaviviruses) can also include in addition to the mutation(s) (e.g., membrane protein mutations) discussed above, one or more mutations in the hinge region or the hydrophobic pocket of the envelope protein, as such mutations have been shown to result in decreased viscerotropism (Monath et al., J. Virol. 76:1932-1943, 2002; WO 03/103571 A2; WO 05/082020; Guirakhoo et al., J. Virol. 78(18):9998-10008, 2004). The polypeptide chain of the envelope protein folds into three distinct domains: a central domain (domain I), a dimerization domain (domain II), and an immunoglobulin-like module domain (domain III). The hinge region is present between domains I and II and, upon exposure to acidic pH, undergoes a conformational change (hence the designation "hinge") that results in the formation of envelope protein trimers that are involved in the fusion of viral and endosomal membranes, after virus uptake by receptor-mediated endocytosis. Prior to the conformational change, the proteins are present in the form of dimers.

Numerous envelope amino acids are present in the hinge region including, for example, amino acids 48-61, 127-131, and 196-283 of yellow fever virus (Rey et al., Nature 375:291-298, 1995). Any of these amino acids, or closely surrounding amino acids (and corresponding amino acids in other Flavivirus envelope proteins), can be mutated according to the invention, and tested for attenuation. Of particular interest are amino acids within the hydrophobic pocket of the hinge region. As a specific example, it has been shown that substituting envelope protein amino acid 204 (K to R), which is in the hydrophobic pocket of the hinge region, in a chimeric Flavivirus including dengue 1 envelope protein sequences inserted into a yellow fever virus vector results in attenuation (Guirakhoo et al., J. Virol. 78:9998-10008, 2004). This substitution leads to an alteration in the structure of the envelope protein, such that intermolecular hydrogen bonding between one envelope monomer and another in the

wild type protein is disrupted and replaced with new intramolecular interactions within monomers. This observation led to a proposal that the attenuation resulting from this substitution is due to these new interactions, which change the structure of the protein in the pre-fusion conformation, most likely by altering the pH threshold that is required for fusion of viral membrane with the host cell, and provides a basis for the design of further attenuated mutants in which additional substitutions are used to increase intramolecular interactions in the hydrophobic pocket, leading to attenuation. Examples of such mutations/substitutions that can be made in the hydrophobic pocket, and included in the viruses of the invention, include substitutions in E202K, E204K, E252V, E253L, E257E, E258G, and E261H (and corresponding substitutions in other Flaviviruses). Any amino acid changes in the corresponding region of the E protein of JE and WN viruses can be designed and incorporated based on the knowledge of homologous protein structure.

affect function and thereby reduce virulence, as described by Hurrelbrink and McMinn (Adv. Virus Dis. 60:1-42, 2003). The functional regions of the E protein in which mutations may be inserted that, together with the membrane deletions/mutations described in the present application, may result in an appropriately attenuated vaccine include: a) the putative receptor binding region on the external surface of domain III, b) the molecular hinge region between domains I and II, which determines the acid-dependent conformational changes of the E protein in the endosome and reduce the efficiency of virus internalization; c) the interface of prM and E proteins, a region of the E protein that interfaces with prM following the rearrangement from dimer to trimer after exposure to low pH in the endosome; d) the tip of the fusion domain of domain II, which is involved in fusion to the membrane of the endosome during internalization events; and e) the stem-anchor region, which is also functionally is involved in conformational changes of the E protein during acid-induced fusion events.

Additional attenuating mutations that can be included with one or more membrane protein mutations in the viruses of the invention include mutations in the 3'untranslated region of the yellow fever virus backbone. The organization of the 3'UTR of a yellow fever virus vaccine strain, YF 17D, which is shared by all ChimeriVax<sup>TM</sup> viruses, is shown in Fig. 1A. It includes in order from the 3' end (i) a 3'-extreme stem-and-loop structure that has been hypothesized to function as a

promoter for minus-strand RNA synthesis and is conserved for all Flaviviruses, (ii) two conserved sequence elements, CS1 and CS2, which share a high degree of nucleotide sequence homology with all mosquito-borne Flaviviruses, and (iii) unique for West African yellow fever virus strains, including the YF17D vaccine virus, three copies of a repeat sequence element (RS) located in the upstream portion of the 3'UTR (Chambers et al., Annu. Rev. Microbiol. 44:649-688, 1990). The 3'UTR also includes numerous stem-loop structures, such as those in the non-conserved region downstream from the RS elements, as depicted in Fig. 1B.

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3'UTR mutations that can be included in the viruses of the invention generally are short, attenuating deletions of, for example, less than 30 nucleotides (e.g., 1, 2, 3, etc., and up to 29 (e.g., 2-25, 3-20, 4-15, 5-10, or 6-8 nucleotides in length); U.S. Patent Application Nos. 60/674,546 and 60/674,415). In some examples, the short 3'UTR deletions are designed to destabilize the secondary structure of one or more of the stem structures in the 3'UTR. In addition to deletions, mutations in such structures can also include substitutions that similarly result in stem structure destabilization. In certain examples, the stem-loop structures that are subject to the mutations are present in nonconserved regions of the 3'UTR or in conserved regions that can tolerate such mutations (e.g., in CS2). For example, the stem destabilizing mutations can be present in any one or more of the predicted stem structures shown in Fig. 1B, which shows four examples of such deletions (dA, dB, dC, and dD). Thus, in addition to these specific examples, other examples of 3'UTR mutations in yellow fever virus include mutations that comprise, e.g., 1-2, 3-8, 4-7, or 5-6 nucleotides of the following stem sequences, which are shown in Fig. 1B as read from 5' to 3': TGGAG, CTCCA, GACAG, TTGTC, AGTTT, GGCTG, CAGCC, AACCTGG, TTCTGGG, CTACCACC, GGTGGTAG, GGGGTCT, AGACCCT, AGTGG, and TTGACG. These mutations can also be present in corresponding amino acids of other Flaviviruses, as described herein.

In addition to stem destabilizing mutations, other short deletions in the 3'UTR can also be included with one or more membrane (and possibly other) mutations in the viruses of the invention. For example, the previously described  $\Delta 30$  mutation (Men et al., J. Virol. 70:3930-3937, 1996; U.S. Patent No. 6,184,024 B1) or mutations that fall within this sequence can be used. Thus, for example, the invention includes any viable deletions that are 1, 2, 3, etc., and up to 29 (e.g., 1-25, 2-20, 3-15, 4-14, 5-13, 6-12, 7-11, 8-10, or 9) nucleotides in length within this region. As a specific example, viruses of the

invention can include deletion d7, in which the following nucleotides from this region in YF17D are deleted: nucleotides 345-351 (AAGACGG; numbering from the 1st nucleotide of the 3'UTR, after the UGA termination codon of the viral ORF; Fig. 1A). Mutations that include deletion of, for example, 1, 2, 3, 4, or 5 additional nucleotides from the 3' or 5' end of this sequence are also included in the invention. In other examples, short deletions in conserved sequences CS1 and CS2 are included in the invention. These mutations can include deletion of, e.g., 1-29, 2-25, 3-20, 4-15, 5-10, or 6-8 nucleotides of these sequences. As two specific examples, nucleotides 360-364 (GGTTA; CS2d5; Fig. 1A) and/or nucleotides 360-375 (GGTTAGAGGAGACCCT; CS2d16; Fig. 1A) are deleted from CS2 of the YF17D-specific 3'UTR. Mutations that include deletion of, for example, 1, 2, 3, 4, or 5 additional nucleotides from the 3' or 5' end of this sequence can also be used. For other flavivirus 3'UTRs, similar mutations can be made, based on the secondary structures of the 3'UTR's. Predictions of secondary structures of 3'UTR of other flaviviruses have been published, e.g., for dengue, Kunjin, and TBE (see, e.g., Proutski et al., Virus Res. 64:107-123, 1999) and HCV (see, e.g., Kolykhalov et al., J. Virol. 70:3363-3371, 1996). Further, numerous 3'UTR nucleotide sequences for many strains of flaviviruses representing all four major serocomplexes (YF, JE, dengue, and TBE) are available from GenBank. Sequences of additional strains can be determined by virus sequencing. The secondary structures of these sequences can be easily predicted using standard software (e.g., mfold or RNAfold programs) to reveal potential stem-loop structures that can be subject to mutagenesis.

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It should be noted that the true secondary structures of the 3'UTRs of Flaviviruses, including YF 17D virus, are unknown because there are no available methods to experimentally prove their existence in the context of whole viruses, and therefore published predictions, e.g., the one predicted for YF 17D by Proutski and coworkers (Fig. 1B), may be incorrect. Many alternative structures can be predicted to form in a relatively long RNA molecule (Zuker et al., N.A.R. 19:2707-2714, 2001), and it is possible that different structures (in plus or minus strands) form and function at different steps of the viral life cycle. True structures can be influenced by the formation of various pseudoknots (Olsthoorn et al., RNA 7:1370-1377, 2001) and long range RNA interactions (e.g., RNA cyclization and other interactions (Alvarez et al., J. Virol. 79:6631-6643, 2005)), as well as possible RNA interactions with host and viral proteins. To further complicate interpretation of published results of theoretical computer predictions, manual

operations are often used, such as initial folding of partial sequences with subsequent forcing of initially predicted structures into structures of longer RNA sequences, the artificial use of N's during initial folding steps, and subjective selection of preferred structure elements (e.g., Mutebi et al., J. Virol. 78:9652-9665, 2004). To this end, we folded the 3'UTR RNA sequence of YF 17D using the commonly used Zuker's prediction algorithm. The predicted optimal structure is shown in Fig. 1C, which differs from the Proutsky prediction shown in Fig. 1B. It is important that the small deletions dA, dB, dC, dD, d7, and d14 in Figs. 1A and 1B generally destabilized the predicted native YF 17D optimal (Fig. 1C) and suboptimal structures. An example of one such altered optimal structure (for the dC mutant) is shown in Fig. 1D. In contrast, the CS2d5 and CS2d16 deletions (Figs. 1A and 1B) did not noticeably change the optimal native structure, indicating that these deletions may attenuate the virus (attenuation was demonstrated in the hamster model for ChimeriVax<sup>TM</sup>-WN) by virtue of altering the sequence of CS2 per se rather than the 3'UTR structure, or alternatively by altering some suboptimal structures. Thus, even though some of the deletions were designed based on the Proutski structure prediction (Fig. 1B), their true effect may be due to destabilizing different structure elements than the predicted stem-loops in Fig. 1B.

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Additional mutations that can be included with membrane protein (and possibly other) mutations in the viruses of the invention are short deletion (e.g., deletions of 1, 2, 3, or 4 amino acids) mutations within the capsid protein. Examples of such mutations, provided in reference to the YF 17D virus capsid protein, include viable deletions affecting Helix I of the protein (see Fig. 2A). A specific example of such a mutation is mutation C2, which includes a deletion of amino acids PSR from Helix I (Fig. 2A). Other short mutations in this region (as well as corresponding mutations in other Flavivirus sequences) can be tested for viability and attenuation, and can also be used in the invention. Capsid protein sequences of other flaviviruses have been published, e.g., for TBE, WN, Kunjin, JE, and dengue viruses (e.g., Pletnev et al., Virology 174:250-263, 1990).

The following are specific examples of chimeric Flaviviruses, which were deposited with the American Type Culture Collection (ATCC) in Manassas, Virginia, U.S.A. under the terms of the Budapest Treaty and granted a deposit date of January 6, 1998, that can be used to make viruses of the invention: Chimeric Yellow Fever

17D/Dengue Type 2 Virus (YF/DEN-2; ATCC accession number ATCC VR-2593) and Chimeric Yellow Fever 17D/Japanese Encephalitis SA14-14-2 Virus (YF/JE A1.3; ATCC accession number ATCC VR-2594). Details of making chimeric viruses that can be used in the invention are provided, for example, in U.S. Patent No.
5 6,696,281 B1; international applications PCT/US98/03894 (WO 98/37911) and PCT/US00/32821 (WO 01/39802); and Chambers et al., J. Virol. 73:3095-3101, 1999, and are also provided below. These methods can be modified for use in the present invention by including a step of introducing one or more mutations as described herein into inserted sequences (e.g., Japanese encephalitis virus or West Nile virus membrane protein or other sequences). Methods that can be used for producing viruses in the invention are also described in PCT/US03/01319 (WO 03/060088 A2; also see below).

Mutations can be made in the viruses of the invention using standard methods, such as site-directed mutagenesis. One example of the type of mutation present in the viruses of the invention is substitutions, but other types of mutations, such as deletions and insertions, can be used as well. In addition, as is noted above, the mutations can be present singly or in the context of one or more additional mutations, whether within the membrane protein itself or in any combination of, e.g., 3'UTR, capsid, or envelope sequences.

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The viruses (including chimeras) of the present invention can be made using standard methods in the art. For example, an RNA molecule corresponding to the genome of a virus can be introduced into primary cells, chick embryos, or diploid cell lines, from which (or the supernatants of which) progeny virus can then be purified. Another method that can be used to produce the viruses employs heteroploid cells, such as Vero cells (Yasumura et al., Nihon Rinsho 21:1201-1215, 1963). In this method, a nucleic acid molecule (e.g., an RNA molecule) corresponding to the genome of a virus is introduced into the heteroploid cells, virus is harvested from the medium in which the cells have been cultured, and harvested virus is treated with a nuclease (e.g., an endonuclease that degrades both DNA and RNA, such as Benzonase<sup>TM</sup>; U.S. Patent No. 5,173,418). In the case of Benzonase<sup>TM</sup>, 15 units/mL can be used, and the conditioned medium refrigerated at 2-8°C for about 16 or more hours to allow for digestion of nucleic acids. The nuclease-treated virus is then concentrated (e.g., by use of ultrafiltration using a filter having a molecular weight

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cut-off of, e.g., 500 kDa (e.g., a Pellicon-2 Mini unltrafilter cassette)), diafiltered against MEME without phenol red or FBS, formulated by the addition of lactose, and filtered into a sterile container. Details of this method are provided in WO 03/060088 A2. Further, cells used for propagation of viruses of the invention can be grown in serum free medium, as described below.

The viruses of the invention can be administered as primary prophylactic agents in those at risk of infection, or can be used as secondary agents for treating infected patients. Because the viruses are attenuated, they are particularly well-suited for administration to "at risk individuals" such as the elderly, children, or HIV infected persons. The vaccines can also be used in veterinary contexts, e.g., in the vaccination of horses against West Nile virus infection, or in the vaccination of domestic pets (e.g., cats, dogs, and birds), livestock (e.g., sheep, cattle, pigs, birds, and goats), and valuable animals such as rare birds. Further, the vaccines of the invention can include a virus, such as a chimeric virus, including a particular mutation (e.g., the M5, M60, and/or M66 mutation), in a mixture with viruses lacking such mutations.

Formulation of the viruses of the invention can be carried out using methods that are standard in the art. Numerous pharmaceutically acceptable solutions for use in vaccine preparation are well known and can readily be adapted for use in the present invention by those of skill in this art (see, e.g., Remington's Pharmaceutical Sciences (18<sup>th</sup> edition), ed. A. Gennaro, 1990, Mack Publishing Co., Easton, PA). In two specific examples, the viruses are formulated in Minimum Essential Medium Earle's Salt (MEME) containing 7.5% lactose and 2.5% human serum albumin or MEME containing 10% sorbitol. However, the viruses can simply be diluted in a physiologically acceptable solution, such as sterile saline or sterile buffered saline. In another example, the viruses can be administered and formulated, for example, in the same manner as the yellow fever 17D vaccine, e.g., as a clarified suspension of infected chicken embryo tissue, or a fluid harvested from cell cultures infected with the chimeric yellow fever virus.

The vaccines of the invention can be administered using methods that are well known in the art, and appropriate amounts of the vaccines to be administered can readily be determined by those of skill in the art. What is determined to be an

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appropriate amount of virus to administer can be determined by consideration of factors such as, e.g., the size and general health of the subject to whom the virus is to be administered. For example, the viruses of the invention can be formulated as sterile aqueous solutions containing between 10<sup>2</sup> and 10<sup>8</sup>, e.g., 10<sup>3</sup> to 10<sup>7</sup> or 10<sup>4</sup> to 10<sup>6</sup>, infectious units (e.g., plaque-forming units or tissue culture infectious doses) in a dose volume of 0.1 to 1.0 ml, to be administered by, for example, intramuscular, subcutaneous, or intradermal routes. In addition, because Flaviviruses may be capable of infecting the human host *via* mucosal routes, such as the oral route (Gresikova et al., "Tick-borne Encephalitis," In *The Arboviruses, Ecology and Epidemiology*, Monath (ed.), CRC Press, Boca Raton, Florida, 1988, Volume IV, 177-203), the viruses can be administered by mucosal (e.g., oral) routes as well. Further, the vaccines of the invention can be administered in a single dose or, optionally, administration can involve the use of a priming dose followed by one or more booster doses that are administered, e.g., 2-6 months later, as determined to be appropriate by those of skill in the art.

Optionally, adjuvants that are known to those skilled in the art can be used in the administration of the viruses of the invention. Adjuvants that can be used to enhance the immunogenicity of the viruses include, for example, liposomal formulations, synthetic adjuvants, such as (e.g., QS21), muramyl dipeptide, monophosphoryl lipid A, or polyphosphazine. Although these adjuvants are typically 20 used to enhance immune responses to inactivated vaccines, they can also be used with live vaccines. In the case of a virus delivered via a mucosal route, for example, orally, mucosal adjuvants such as the heat-labile toxin of E. coli (LT) or mutant derivations of LT can be used as adjuvants. In addition, genes encoding cytokines that have adjuvant activities can be inserted into the viruses. Thus, genes encoding 25 cytokines, such as GM-CSF, IL-2, IL-12, IL-13, or IL-5, can be inserted together with foreign antigen genes to produce a vaccine that results in enhanced immune responses, or to modulate immunity directed more specifically towards cellular, humoral, or mucosal responses. Additional adjuvants that can optionally be used in 30 the invention include toll-like receptor (TLR) modulators.

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In the case of dengue viruses and/or chimeric Flaviviruses including membrane and envelope proteins of a dengue virus, against which optimal vaccination can involve the induction of immunity against all four of the dengue serotypes, the viruses of the invention can be used in the formulation of tetravalent vaccines. Any or all of the viruses used in such tetravalent formulations can include one or more mutations that decrease viscerotropism, as is described herein. The viruses can be mixed to form tetravalent preparations at any point during formulation, or can be administered in series. In the case of a tetravalent vaccine, equivalent amounts of each virus may be used. Alternatively, the amounts of each of the different viruses present in the administered vaccines can vary (WO 03/101397 A2).

The invention also includes nucleic acid molecules (e.g., RNA or DNA (e.g., cDNA) molecules) that correspond to the genomes of the viruses of the invention as described herein, or the complements thereof. These nucleic acid molecules can be used, for example, in methods of manufacturing the viruses of the invention. In such methods, a nucleic acid molecule corresponding to the genome of a virus is introduced into cells in which the virus can be produced and replicate (e.g., primary cells, chick embryos, diploid cell lines, or heteroploid cell lines (e.g., Vero cells)), and from which (or the supernatants of which) progeny virus can then be purified. These methods can further include virus purification steps, as is known in the art.

As is noted above, details of making chimeric viruses that can be used in the invention are provided, for example, in U.S. Patent No. 6,696,281 B1; international applications PCT/US98/03894 (WO 98/37911) and PCT/US00/32821 (WO 01/39802); and Chambers et al., J. Virol. 73:3095-3101, 1999. Details of the construction of a chimeric Flavivirus including pre-membrane and envelope proteins of Japanese encephalitis virus (or West Nile virus), and capsid and non-structural proteins of yellow fever virus, are provided as follows. These methods can readily be adapted by those of skill in the art for use in constructing chimeras including the mutations described herein, as well as chimeras including other pre-membrane and envelope sequences.

Briefly, derivation of a YF/JE chimera can involve the following. YF genomic sequences are propagated in two plasmids (YF5'3'IV and YFM5.2), which encode the YF sequences from nucleotides 1-2,276 and 8,279-10,861 (YF5'3'IV) and

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from 1,373-8,704 (YFM5.2) (Rice et al., The New Biologist 1:285-296, 1989). Full-length cDNA templates are generated by ligation of appropriate restriction fragments derived from these plasmids. YF sequences within the YF5'3'IV and YFM5.2 plasmids are then replaced by the corresponding JE sequences from the start of the prM protein (nucleotide 478, amino acid 128) through the E/NS1 cleavage site (nucleotide 2,452, amino acid 817).

RNA was extracted from monolayers of infected LLC-MK2 cells and first strand synthesis of negative sense cDNA was carried out using reverse transcriptase with a negative sense primer (JE nucleotide sequence 2,456-71) containing nested XbaI and NarI restriction sites for cloning initially into pBluescript II KS(+), and subsequently into YFM5.2(NarI), respectively. First strand cDNA synthesis was followed by PCR amplification of the JE sequence from nucleotides 1,108-2,471 using the same negative sense primer and a positive sense primer (JE nucleotides sequence 1,108-1,130) containing nested XbaI and NsiI restriction sites for cloning into pBluescript and YFM5.2(NarI), respectively. JE sequences were verified by restriction enzyme digestion and nucleotide sequencing. The JE nucleotide sequence from nucleotides 1 to 1,130 was derived by PCR amplification of negative strand JE cDNA using a negative sense primer corresponding to JE nucleotides 1,116 to 1,130 and a positive sense primer corresponding to JE nucleotides 1 to 18, both containing an EcoRI restriction site. PCR fragments were cloned into pBluescript and JE sequences were verified by nucleotide sequencing. Together, this represents cloning of the JE sequence from nucleotides 1-2,471 (amino acids 1-792).

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To insert the C terminus of the JE envelope protein at the YF E/NS1 cleavage site, a unique NarI restriction site was introduced into the YFM5.2 plasmid by oligonucleotide-directed mutagenesis of the signalase sequence at the E/NS1 cleavage site (YF nucleotides 2,447-2,452, amino acids 816-817) to create YFM5.2(NarI).

Transcripts derived from templates incorporating this change were checked for infectivity and yielded a specific infectivity similar to the parental templates (approximately 100 plaque-forming units/250 nanograms of transcript). The JE sequence from nucleotides 1,108 to 2,471 was subcloned from several independent PCR-derived clones of pBluescript/JE into YFM5.2(NarI) using the unique NsiI and NarI restriction sites. YF5'3'IV/JE clones containing the YF 5' untranslated region (nucleotides 1-118) adjacent to the JE prM-E region were derived by PCR amplification.

To derive sequences containing the junction of the YF capsid and JE prM, a negative sense chimeric primer spanning this region was used with a positive sense primer corresponding to YF5'3'IV nucleotides 6,625-6,639 to generate PCR fragments that were then used as negative sense PCR primers in conjunction with a positive sense primer complementary to the pBluescript vector sequence upstream of the *EcoRI* site, to amplify the JE sequence (encoded in reverse orientation in the pBluescript vector) from nucleotide 477 (N-terminus of the prM protein) through the *NheI* site at nucleotide 1,125. The resulting PCR fragments were inserted into the YF5'3'IV plasmid using the *NotI* and *EcoRI* restriction sites. This construct contains the SP6 promoter preceding the YF 5'-untranslated region, followed by the sequence: YF (C) JE (prM-E), and contains the *NheI* site (JE nucleotide 1,125) required for *in vitro* ligation.

To use the *NheI* site within the JE envelope sequence as a 5' in vitro ligation site, a redundant *NheI* site in the YFM5.2 plasmid (nucleotide 5,459) was eliminated. This was accomplished by silent mutation of the YF sequence at nucleotide 5,461 (T C; alanine, amino acid 1820). This site was incorporated into YFM5.2 by ligation of appropriate restriction fragments and introduced into YFM5.2(*NarI*)/JE by exchange of an *NsiI/NarI* fragment encoding the chimeric YF/JE sequence.

To create a unique 3' restriction site for *in vitro* ligation, a *BspE*I site was engineered downstream of the *Aat*II site normally used to generate full-length templates from YF5'3'IV and YFM5.2. (Multiple *Aat*II sites are present in the JE

structural sequence, precluding use of this site for *in vitro* ligation.) The *BspE*I site was created by silent mutation of YF nucleotide 8,581 (A C; serine, amino acid 2,860), and was introduced into YFM5.2 by exchange of appropriate restriction fragments. The unique site was incorporated into YFM5.2/JE by exchange of the *Xbal/Sph*I fragment, and into the YF5'3'IV/JE(prM-E) plasmids by three-piece ligation of appropriate restriction fragments from these parent plasmids and from a derivative of YFM5.2 (*BspE*I) deleting the YF sequence between the *EcoR*I sites at nucleotides 1 and 6,912.

cDNA from a clone of the JE Nakayama strain, which has been extensively characterized in expression experiments and for its capacity to induce protective immunity (see, e.g., McIda et al., Virology 158:348-360, 1987; the JE Nakayama strain is available from the Centers for Disease Control, Fort Collins, Colorado, and the Yale Arbovirus Research Unit, Yale University, New Haven, Connecticut), was also used in the construction of chimeric flaviviruses. The Nakayama cDNA was inserted into the YF/JE chimeric plasmids using available restriction sites (*Hind*III to *Pvu*II and *Bpm*I to *Mun*I) to replace the entire prM-E region in the two plasmid system except for a single amino acid, serine, at position 49, which was left intact in order to utilize the *Nhe*I site for *in vitro* ligation.

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Procedures for generating full-length cDNA templates are essentially as described in Rice et al. (The New Biologist 1:285-96, 1989). In the case of chimeric templates, the plasmids YF5'3'TV/JE (prM-E) and YFM5.2/JE are digested with Nhel/BspEI and in vitro ligation is performed using 300 nanograms of purified fragments in the presence of T4 DNA ligase. The ligation products are linearized with XhoI to allow run-off transcription. SP6 transcripts are synthesized using 50 nanograms of purified template, quantitated by incorporation of <sup>3</sup>H-UTP, and integrity of the RNA is verified by non-denaturing agarose gel electrophoresis. Yields range from 5 to 10 micrograms of RNA per reaction using this procedure, most of which is present as full-length transcripts. Transfection of RNA transcripts in the presence of cationic liposomes is carried out as described by Rice et al. (supra) for YF 17D, to generate the chimeric viruses.

In the case of chimeric flaviviruses including West Nile virus and yellow fever virus sequences, the two-plasmid system described above can also be used. In one example, the West Nile (WN) virus prM and E genes used were cloned from WNV

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flamingo isolate 383-99, sequence GenBank accession number AF196835. Virus prME cDNA was obtained by RT-PCR (XL-PCR Kit, Perkin Elmer). The 5' end of WN prM gene was cloned precisely at the 3'end of the YF 17D capsid gene by overlap-extension PCR using Pwo polymerase (Roche). The 3' end of the E gene was also cloned precisely at the 5'end of the YF NS1 coding sequence by overlap-extension PCR. Silent mutations were introduced into the sequence of prM and E to create unique restriction sites Bsp EI and Eag I. Digestion of the two plasmids with these enzymes generated DNA fragments that were gel purified and ligated in vitro to produce a full-length chimeric cDNA. The cDNA was linearized with Xho I to facilitate in vitro transcription by SP6 polymerase (Epicentre). The RNA product was introduced into eukaryotic cell lines permissive for viral RNA translation and replication of the virus. As with the YF/JE chimera, described above, mutations of the invention can be introduced into YF/WN chimeras as described herein, using standard methods.

Other Flavivirus chimeras can be engineered with a similar strategy, using natural or engineered restriction sites and, for example, oligonucleotide primers as shown in Table 14.

The invention is based, in part, on the experimental results described in the following Examples.

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#### **EXAMPLES**

### Example 1: ChimeriVax<sup>TM</sup>-WN

#### **Experimental Results**

Background and Summary

A chimeric yellow fever-West Nile (YF-WN) virus, ChimeriVax<sup>TM</sup>-WN, was produced by insertion of pre-membrane (prM) and envelope (E) genes of a WN virus (NY99) into the YF17D backbone. The virus was produced in Vero cells under serum free conditions (at Passage 5, P5), evaluated for safety, immunogenicity, and efficacy in preclinical models, and has been tested in a phase I study in humans. Additional attenuation of the vaccine virus (P5) is determined by three SA14-14-2-specific mutations in the E protein (residues 107, 316, and 440). The vaccine virus was less neurovirulent than YF-VAX<sup>®</sup> when tested in mouse and monkeys inoculated by the IC route and protected mice, hamsters, and monkeys upon a single inoculation

(Arroyo et al., J. Virol. 78:12497-12507, 2004; Tesh et al., Emer. Infect. Dis. 8:1392-1397, 2002). The vaccine virus contained a mixed population of viruses (exhibiting small, S, and large, L, plaque phenotypes), which differed by a single amino acid residue in the M protein at position 66 (M66). This mutation did not affect
5 neurovirulence of the virus for 8 day old suckling mice (Arroyo et al., J. Virol. 78:12497-12507, 2004). In the current invention, we describe the discovery that the M66 mutation reduces viremia in the host and thus can be used to improve the safety of the current vaccine (ChimeriVax<sup>TM</sup>-WN02, P5, mixed population of parent and mutant viruses) or the large plaque variant (non mutant) virus.

A nucleotide heterogeneity (~50%) of T and C (CTA/CCA) was observed in the consensus sequence of ChimeriVax<sup>TM</sup>-West Nile vaccine virus at P5 produced in Vero cells under serum free conditions. This mutation would result in presence of viruses containing either amino acid Proline (mutant) or Leucine (parent wild type) in the membrane (M) protein at residue 66 (herein designated as M66 mutation). The sequences of ChimeriVax<sup>TM</sup>WN02 and the ChimeriVax<sup>TM</sup>WN02 M66 variant are provided in the enclosed sequence appendix, which also includes an alignment of the amino acid sequences of these proteins.

The M protein of West Nile virus contains 75 amino acids. The structure of

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the protein was predicted and compared to the structures of M proteins of JE SA14 20 (AAA67174), Kunjin (AAP78942), MVE (CAA27184), SLE MSI (AAP44973, and SLE CORAN (AAP44972) by submission of the protein sequence to the http://www.predictprotein.org website. In all predicted structures, the first 40 amino acids of the M protein (SLTVQTHGESTLANKKGAWMDSTKATRYLVKTESWILRN) are predicted to be a non-membrane region, whereas the remaining 35 amino acids (40-75) 25 (PGYALVAAVIGWMLGSNTMQRVVFVVLLLLVAPAYS) are predicted to be within the viral membrane region. In addition, there are 9-10 charged amino acids (3-4 acidic, E or D) and 6 basic (R or K) within the first 40 amino acid residues, whereas there is only one charged amino acid (basic) at residue 60 of all 5 Flaviviruses (WNV, SLE, MVE, JE, and Kunjin) described here. Thus, it may be that the M60 residue 30 plays a vital role in biology of the virus by interaction within its neighboring amino acids.

The plaque morphology of the vaccine virus at P5 revealed a mixed population of L and S plaque size phenotypes. The sequencing of the P2, P3, P4, and P5 viruses revealed that the mutation first appeared at P4 (10% of the total population) and reached ~50% in P5. The sequencing of the S and L plaque isolates of the vaccine virus showed that the mutation is responsible for a change in plaque size from L to S. Both S and L virus variants (prepared as research virus) did not significantly differ in their neurovirulence for 8 day old suckling mice (p=<0.0001).

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Pre-Master Seed (PMS, P10) stocks of both L and S viruses were produced in Vero cells from ChimeriVax<sup>TM</sup>-WN02 (p5) under "clean laboratory condition" by 3 rounds of direct plaque to plaque purifications and 2 rounds of virus amplification. 10 The sequencing of P10 S and L viruses revealed a single amino acid difference in the M66 residue (S virus contained Proline at M66 residue, whereas L virus contained Leucine at this site). The M66 mutation seemed to be stable under large scale manufacturing conditions. When the S plaque virus (P10, PMS) was inoculated into hamsters by subcutaneous inoculation, it induced a very low level of viremia 15 compared to the vaccine virus (P5) or the L plaque virus variant (P10, PMS). In sera of monkeys and humans inoculated with ChimeriVax<sup>TM</sup>-WN P5 virus (contained ~50:50 S and L plaque variants), the majority of viruses were of L plaque size phenotype. In addition, it was shown that the S plaques grow to a lower titer than the L plaques in human hepatoma cell lines. These data indicated that the S plaque virus 20 (ChimeriVax<sup>TM</sup>-WN02 with M66 mutation) may induce a lower level of viremia in humans than ChimeriVax<sup>TM</sup>-WN02 (without M66 mutation), and therefore could constitute a suitable (safe) WN vaccine candidate for "at risks individuals," such as the elderly, children, or HIV infected persons. Additional mutations in the M region were found by sequencing individual plaques isolated from large scale manufacturing 25 passages (e.g., M62, M63, and M64) of PMS S plaque from P10 to P12 or monkeys inoculated with ChimeriVax<sup>TM</sup>-WN02 vaccine (e.g., M60, M61, and M63). These mutations can also be used in the construction of viruses of the invention.

30 Production of PMS of S and L plaque viruses in Vero cells

ChimeriVax<sup>TM</sup>-WN02 vaccine material (P5) was grown in Serum Free Vero cells; 10 plaques identified as "small" (S) and 10 plaques identified as "large" (L) were picked. Each isolate was then passaged on Serum Free Vero cells, and one

plaque was picked from each isolate. The procedure was repeated one final time, for a total of three rounds of plaque purification. The plaque purified isolates (P8) were amplified in T25 cm<sup>2</sup> flasks containing Serum Free Vero cells (and grown in serum free (SF) media), then harvested and stored at -80°C. Isolates were sequenced to find a PMS candidate free of spurious mutations. Two isolates were identified to be free of expressed (non-silent) mutations: one isolate was confirmed to be small plaque (M66 Proline) (Table 1), and the other contained a wt sequence (M66 Leucine) (Table 2). These two isolates were then grown in large flasks, aliquoted, and submitted to QC inventory as LP and SP PMS (P10) viruses.

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Genetic stability of SP viruses produced at large scale

In order to determine if the S plaque phenotype is stable during a large scale manufacturing process, the small plaque PMS virus was passaged twice in a bioreactor by infecting Vero cells and growing under serum free conditions to produce the P12 virus. The P12 virus was harvested and plaqued in 6-well plates. The majority of the plaques were of small size. Twenty of the largest plaques available were picked, amplified on O-Vero (one passage), and the prME region was transcribed/amplified via Titan One-Tube RT-PCR kit (Roche). The cDNA fragments containing the M region were sequenced, and the morphology of the isolates was confirmed via immuno-staining using WN specific monoclonal antibodies. Thirteen of 20 plaques contained only M66 (the genetic marker responsible for SP morphology), and 5 isolates contained other mutations in addition to M66. Isolate #4 contained M63 (LP phenotype), and isolate #16 contained a mixed population of wt and M66. These data demonstrated that, despite the fact that some plaques appeared to be of large size, they contained the M66 mutation and upon amplification proved to be of S size. Only one plaque (#4) out of 20 appeared to be of L size, apparently due to a mutation at M63 from L to P. Plaque #16 appeared to produce a mixed population of large and small plaque size viruses containing both wt L and mutant P amino acids at position M66 (Table 3).

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Growth of ChimeriVax<sup>TM</sup>-WN virus variants in hepatic cells

Human hepatoma cell lines HepG2 and THLE-3 cells were infected with ChimeriVax<sup>TM</sup>-WN01 (wild type prME), ChimeriVax<sup>TM</sup>-WN02 P5 (containing

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mutations at E107, E313, E316, E440, M66 mixed L/P amino acids, mixed S and L plaques), ChimeriVax<sup>TM</sup>-WN LP (E107, E313, E316, and E440, WNL), and ChimeriVax<sup>TM</sup>-WN SP (E107, E313, E316, E440, and M66P, WNS) at an MOI of 0.005. Supernatants were collected daily and titrated on O-Vero cells using the standard neutral red double agarose overlay procedure.

In HepG2 cells (Fig. 3) the highest virus growth (7x10<sup>6</sup> PFU/ml) was observed on Day 5 with WN01 (wild type prME), followed by that of LP (2.7x10<sup>6</sup> PFU/ml) on Day 5. The virus peak with YF-VAX® was reached on Day 3 (1.17x10<sup>6</sup> PFU/ml), followed by WN02 mixed vaccine virus (6.4x10<sup>5</sup> PFU/ml) on Day 4. The lowest growth was found with the SP virus (peak titer on Day 4 was 6.1x10<sup>5</sup> PFU/ml), which contained a single amino acid substitution (L to P) at M66. In THLE-3 cells (Fig. 4), the same pattern as in HepG2 cells was observed with the exception that the titer of YF-VAX® was slightly higher than that of the LP virus. The highest titer was seen again with the WN01 (1.3x10<sup>5</sup> PFU/ml, Day 4), followed by those of LP (5.7x10<sup>4</sup> PFU/ml, Day 7), YF-VAX® (8.8 x10<sup>4</sup> PFU/ml, Day 4), and the mixed P5 virus (1.8x10<sup>4</sup> PFU/ml, Day 4). The lowest titer was observed again with the SP virus (9.2x10<sup>3</sup> PFU/ml, Day 4).

The induction of cytopathic effects (CPE) was recorded daily for each virus (Table 4). The CPE for WN 01 and the LP virus was first observed on Day 5 and was completed (100%) 2 days later, whereas SP or mixed plaque population induced CPE at an earlier time point (Day 3) and completely destroyed the cell monolayer one day earlier (Day 6) than WN01 or the LP. The induction of CPE with YF-VAX® was first observed on Day 3 and the monolayer was fully destroyed by Day 6 post inoculation. The induction of CPE in HepG2 cells may be due to apoptotic activity of the M protein, as has been shown with wild type dengue viruses (Catteau et al., J. Gen. Virol. 84:2781-2793, 2003). These data showed that the SP virus variant grows to a lower titer than those of mixed or LP viruses, indicating that the M66 mutation may have rendered the virus less hepatotropic for humans.

Lack of detection of ChimeriVax<sup>TM</sup>-WN, SP viruses after inoculation of monkeys with mixed (SP and LP viruses) P5 vaccine virus

A total of 8 naïve cynomolgus monkeys that lacked detectable antibodies to Flaviviruses, such as WN, JE, and YF viruses (as determined by plaque reduction

neutralization test (PRNT)), were inoculated by the subcutaneous route with either ChimeriVax<sup>TM</sup>-WN02 (P5) (n=4) or YF-VAX<sup>®</sup> (n=4). The purpose of this study was to evaluate viremia, biodistribution, and possible toxicity of the ChimeriVax<sup>TM</sup>-WN02 vaccine during a 3 day observation period. The inoculated dose was ~1.25x10<sup>5</sup> PFU/0.5 mL and 5.5x10<sup>4</sup> PFU/mL for ChimeriVax<sup>TM</sup>-WN02 and YF-VAX<sup>®</sup>, respectively. Animals were bled daily and sacrificed on Day 4 post inoculation. Blood was used to determine the viremia level using a standard plaque assay on Vero cells, whereas collected tissues were either flash frozen for viral analysis or preserved for histopathological evaluations.

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Viremia was assessed on monkey sera collected from Day 1 (before inoculation) through Day 4 (prior to euthanization). The assay was performed either by using agarose double overlay and neutral red staining (to isolate and sequence individual plaques) or by methyl cellulose overlay and crystal violet staining (to measure the level of viremia) as described (Monath et al., J. Virol. 74(4):1742-1751, 2000). The magnitude and duration of viremia in ChimeriVax<sup>TM</sup>-WN02 inoculated monkeys were higher than those of YF-VAX<sup>®</sup> (Table 5). The highest titer of viremia for YF-VAX<sup>®</sup> was 200 PFU/mL (animal MF21157, Day 4). The highest titer of viremia for ChimeriVax<sup>TM</sup>-WN P5 virus was 1000 PFU/mL (animal MF21191F, Day 4). All animals (4/4) inoculated with ChimeriVax<sup>TM</sup>-WN02 virus were viremic for 3 days post inoculation, whereas only 2/4 animals inoculated with YF-VAX<sup>®</sup> became viremic (for only 2 days) (Table 5).

Because animals inoculated with ChimeriVax<sup>TM</sup>-WN02 virus had received a mixture of SP and LP viruses, it was necessary to isolate various SP and LP viruses from sera to identify the virus variant (S or L) responsible for the high level of viremia. Sera of all 4 monkeys obtained from Day 2 to Day 4 post inoculation were diluted 1:2 and 1:10 and used to inoculate duplicate wells of 6-well plates seeded with Vero cells. After addition of the second agarose overlay with neutral red, individual plaques (4 S and 3 L plaques) were picked and directly sequenced to identify the presence of the M66 mutant virus (Table 6). None of the isolated plaques contained the M66 mutation (L to P substitution), indicating that the M66 mutant virus is not responsible for the high level of viremia that was detected in these animals. Interestingly, 3 other mutations were observed in the M region (M60, M61, and M63).

It is possible that either these virus variants had existed in low quantity in the ChimeriVax<sup>TM</sup>-WN02 vaccine virus (which could not be detected by consensus sequencing), or that they have been generated *in vivo* (monkeys) by mutations in the genome of the LP virus variants.

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Viremia and neutralizing antibody responses in hamsters inoculated with ChimeriVax<sup>TM</sup>-WN SP (PMS, P10), LP (PMS, P10), or mixed (P5, SP, and LP) viruses

The animals used in this study were maintained in microseparators under BL2 and handled according to an animal protocol approved by the IACUC throughout the study. Three ChimeriVax<sup>TM</sup>-WN02 viruses (SP, PMS, P10; LP, PMS, P10, and the mix SP and LP vaccine virus, P5) were used to infect 7 week-old female Golden Syrian hamsters (*Mesocricetus auratus*) from Harlan Sprague-Dawley. Each virus was injected into a group of 15 hamsters via the subcutaneous route in the inguinal area. The infection dose was 10<sup>5</sup> pfu, and the inoculum volume was 100 μl. An additional group of 5 animals was similarly injected with 100 μl of virus diluent as sham control. On the day of virus infection (Day 0) and each following day until 5 days post infection, blood samples were collected by retro orbital bleeding from all animals except the sham control group. The animals were anaesthetized by inhalation of isofluorane to effect prior to bleeding and inoculation. Virus concentration in the test samples were determined by direct plaquing of a 0.1 mL of 1:10 diluted serum sample in duplicate wells of Vero cell culture grown in 12-well-plates (Fig. 5).

As is shown in Fig. 5, a higher level (3 logs of pfu on average) of peak viremia was observed in serum samples collected from LP virus infected hamsters, while a very low level (< 10 pfu) of viremia was seen in blood samples of SP virus inoculated hamsters. When the proportion of SP virus was increased (to 50% as for the mixed plaque virus) in the inoculum, the peak viremia titer was lowered to approximately half of the LP virus induced viremia level. Additionally, the viremia peak time was delayed for at least 1 day to 4 days post infection.

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These data demonstrated that the LP and SP variants isolated from the same parent virus, ChimeriVax<sup>TM</sup>-WN02, have different biological properties. The LP virus replicated to a higher level at a faster rate, in comparison with the SP virus in

hamsters. In addition, mixing SP virus with LP (P5 virus) apparently counteracts some properties of the LP virus. This is shown in the hamster infection experiments, in which the presence of virus in blood was reduced to lower levels and the virus replication kinetics were slowed in mixed virus infected hamsters. In sum, the mutation at M66 (L to P) present in SP variant virus significantly reduced its viremia in hamsters.

# Example 2: ChimeriVax<sup>TM</sup>-JE and ChimeriVax<sup>TM</sup>-DEN1-4 Background and Summary

In the study described below, we prepared and characterized a new ChimeriVax<sup>TM</sup>-JE seed virus using Vero cells grown in serum-free (SF) media in order to eliminate any concerns about possible contamination of the vaccine with the prion agent of bovine transmissible encephalopathy. During propagation in SF culture, uncloned virus accumulated mutations not seen previously in serum-containing culture, which appeared to be adaptations to SF growth conditions increasing the rate of virus replication. These mutations occurred in the E or M proteins (E-107 F to L or M-60 R to C mutations) and suggested a functional significance of the M protein in the process of virus replication, which became noticeable during virus growth in SF conditions (see amino acid R at position 60 of the M-protein shown in Example 1 (ChimeriVax<sup>TM</sup>-WN). The effects of mutations within the M (M60, M5 in ChimeriVax<sup>TM</sup>-JE) or the E proteins (E-107 in ChimeriVax<sup>TM</sup>-JE, E202/204 in ChimeriVax<sup>TM</sup>-DEN1 and -DEN3 and E251 in ChimeriVax<sup>TM</sup>-DEN2) on biological properties of the vaccine were defined. All of these chimeric viruses have already been tested in clinical trials.

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#### Materials and methods

Cells and media

Vero cells were originally received from the American Type Culture Collection (ATCC; Manassas, VA; CCL 81; African green monkey kidney cells). These cells were adapted to grow in SF media and were obtained from Baxter (Orth, Austria) at passage 133, and then were used directly by seeding into flasks or seeded

starting from a cell bank at passage 136. In all experiments, the passage level of the Vero cells did not exceed passage 149. Cells and viruses were grown at 36°C under 7.5% CO<sub>2</sub>. Cells were propagated under SF conditions.

## 5 ChimeriVax<sup>TM</sup>-JE variants

The virus was initiated (passage P1) by electroporation of SF Vero cells with the same *in vitro* RNA transcripts (stored at -80°C) that were used previously for production of a non-SF ChimeriVax<sup>TM</sup>-JE vaccine candidate tested in preclinical and clinical trials (Monath et al., Vaccine 20:1004-1018, 2002) and prepared as described previously (Chambers et al., J. Virol. 73:3095-3101, 1999). Amplification passages were generally done at an MOI of 0.001 pfu/cell and viral harvests were collected on days 3-4 postinfection (when CPE was ~ 10%), clarified by slow speed centrifugation, supplemented with 10% sorbitol, and stored at -80°C. Cloned variants were produced in Baxter Vero cells by three consecutive plaque purifications using a standard agarneutral red overlay method in the presence of gamma-irradiated FBS (HyClone; FBS was used because the cells failed to form plaques under agar prepared with SF media) followed by amplification in SF conditions. Plaque assays to determine virus titers in indicated samples were performed using a single methyl cellulose overlay method with visualization of plaques by crystal violet on day 5 post-infection.

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#### ChimeriVax<sup>TM</sup>-DEN viruses

ChimeriVax<sup>TM</sup>-DEN1-4 vaccine viruses were prepared by electroporation of Vero cells with RNA transcripts prepared from viral cDNA. Progeny viruses were subjected to three rounds of plaque purification to produce the Pre-Master Seed (PMS) viruses at passage 7 (P7). Three further passages were carried out using U.S. current Good Manufacturing Practices (cGMP) to produce the Vaccine lot (P10 viruses). Some mutations appeared in the E genes of the chimeras after multiple passages in Vero cells (Guirakhoo et al., J. Virol. 78:4761-4775, 2004). One of these mutations (E 204 in ChimeriVax<sup>TM</sup>-DEN1) significantly reduced viscerotropism of the virus in non-human primates (Guirakhoo et al., J. Virol. 78:9998-10008, 2004).

#### Consensus sequencing

Consensus sequencing of indicated virus samples was performed as previously described (Pugachev et al., Vaccine 20:996-999, 2003). Briefly, virion RNA extracted with the TRIZOL LS reagent (Life Technologies-Gibco BRL) was amplified in five overlapping cDNA amplicons of ~ 2-3 kb in length with Titan One-Tube RT-PCR kit (Roche). Amplicons were sequenced using a collection of JE- and YF-specific oligonucleotide primers of both positive and negative orientation and CEQ Dye Terminator Cycle Sequencing kit (Beckman). Sequencing reaction products were resolved with a CEQ2000XL automated sequencer (Beckman Coulter).

The data were aligned and analyzed with Sequencher 4.1.4 (GeneCodes) software. Nucleotide heterogeneities were registered when a heterogeneous signal was observed in all chromatograms representing both plus- and minus-strand sequencing reactions. For some viruses, only the first of the five cDNA amplicons (Fragment I) was sequenced that includes the structural genes.

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#### Neurovirulence in suckling mice

Institutes of Health guidelines for the humane use of laboratory animals. Pregnant outbred ICR female mice were purchased from Taconic Farms (Germantown, NY).

Newborn mice were fostered and mixed into new groups 6 days prior to inoculation. Groups of 8 day-old suckling mice were inoculated with 0.02 ml of the indicated virus samples by the intracerebral (IC) route. Serial 1:10 dilutions of viruses used for inoculations were done in MEM-10% FBS. Undiluted inocula were back-titrated and the exact doses of each dilution were calculated. Mortalities were recorded over a period of 21 days. The YF 17D control virus was YF-VAX® (Aventis Pasteur, Swiftwater, PA) reconstituted from a commercial vaccine vial.

### Monkey safety and efficacy tests

Experiment 1. The neurovirulence/toxicity profile of new clone C (M-60)

ChimeriVax<sup>TM</sup>-JE Vaccine Master Viral Bank (MVB; P11) and Production Viral

Bank (PVB; P12) stocks, as compared to YF-VAX® control (YF 17D vaccine virus),

was studied according to GLP standards in cynomolgus monkeys. Thirty-three (33)

experimentally-naïve, Flavivirus-seronegative cynomolgus monkeys (as determined

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by HAI test) were assigned to treatment groups as shown in Table 9. All monkeys were dosed via a single IC injection on Day 1, observed for 30 days, and then euthanized and necropsied. The monkeys were evaluated for clinical signs (twice daily), and changes in food consumption (daily), body weight (weekly), and clinical pathology indices. Clinical scores were assigned according to a clinical scoring system, based on the World Health Organization (WHO) requirements for yellow fever vaccine (WHO, Technical Report Series, No. 872, 1998). Blood samples were collected pre-inoculation on Day 1 and on Days 3, 5, 7, 15, and 31 for clinical pathology analysis (serum chemistry and hematology parameters). Additional blood samples were collected on Day 1 (pre-dose) and Days 2-11 for quantitative viremia determinations, and on Day 1 (pre-dose) and Day 31 for neutralizing antibody titer analyses. A complete necropsy was performed on Day 31 and tissues collected for preservation. Tissue was prepared for histopathology of the liver, spleen, heart, kidney, and adrenal glands. Histopathology of the brain and spinal cord was performed according to the methods described by Levenbook et al. (J. Biol. Stand. 15:305, 1987) and incorporated into the WHO requirements for the yellow fever vaccine (WHO, 1998).

Experiment 2. This experiment was conducted to compare the viremia, immune response, and safety of ChimeriVax<sup>TM</sup>-JE Vaccine [original uncloned vaccine P5 produced previously in LS5 Vero cells in the presence of FBS (BB-IND #9167, Serial #000) containing no mutations except for an E491 L to F change in the hydrophobic tail of E protein] and new Clone C (M-60 mutant) ChimeriVax<sup>TM</sup>-JE purified vaccine bulk preparation (P13) over a 30-day period following a single subcutaneous (SC) administration in cynomolgus monkeys according to GLP standards. Eighteen (18) experimentally-naïve, Flavivirus-seronegative (by HAI test) cynomolgus monkeys were assigned to treatment groups as shown in Table 10. All monkeys were dosed once on Day 1 via SC injection at a single site in one arm. The monkeys were evaluated for clinical signs of toxicity (twice daily), changes in body weight (weekly), and serum chemistry, hematology, and coagulation parameters. Blood samples were collected on Day 1 (pre-inoculation) and Days 4, 7, 15, and 31 for serum chemistry, hematology, and coagulation parameter analysis. Additional

blood samples were collected on Day 1 (pre-inoculation) and Days 2-11 for quantitative viremia analysis, and on Day 1 (pre-inoculation) and Day 31 for Japanese encephalitis virus-specific serum antibody titer analysis.

5 pH threshold of virus inactivation (Indirect Fusion Assay)

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One of the consequences of exposure of Flaviviruses to low pH (in the absence of cell membranes) is induction of irreversible conformational changes in the E protein and virus inactivation (loss of potency). In the presence of cell membranes, these conformational changes are necessary for fusion of viral membrane with those of cellular membranes, resulting in release of viral genome into the host cells. The pH threshold for fusion of mosquito-borne viruses such as WN, DEN, YF, and JE can be measured by fusion from within (FFWI) using the mosquito cell line C6/36 (Guirakhoo et al., Virology 169(1):90-99, 1989). We were not, however, able to demonstrate any FFWI with all of our ChimeriVax<sup>TM</sup> viruses, probably due to lack of sufficient growth of these viruses in mosquitoes and mosquito cell lines (Johnson et al., Am. J. Trop. Med. Hyg. 70(1):89-97, 2004). We therefore attempted to measure the loss of virus potency after exposure to different pH levels, in an assay designated here as an "Indirect Fusion Assay." This assay determines indirectly the pH threshold at which the fusion of viral membranes with those of cellular membranes occurs.

Fusion was performed at pH 7.0, 6.8, 6.6, 6.4, 6.2, 6.0, 5.8, 5.6, 5.4, and 5.0, using 1X MEM supplemented with 2 mM L-Glutamine, 2.7% sodium bicarbonate, 10% HI FBS, and 1% antibiotic/antimycotic solution [(100 U/ml of penicillin, 0.1 mg/ml of streptomycin, 0.25 μg/ml Amphotericin (Sigma)] adjusted to the proper pH with MES (Sigma). An aliquot of each virus at 1x10<sup>4</sup> plaque forming unit (PFU)/ml was diluted (10<sup>-1</sup> dilution) in each pH medium. After 10 minutes of exposure at each pH value, 50% heat inactivated (HI) FBS was added to each vial and the pH of each solution was neutralized with sodium bicarbonate. A volume of 100 μl of each virus at each pH value was used to infect Vero-cell monolayers (seeded at a density of 9x10<sup>5</sup> cells/well, in 6-well plates) to determine its titer. Infection was performed in duplicate, so as to cause 50 PFU/well; two non-infected wells of cells were kept per plate and served as negative controls. The pH 7.0 and 6.8 samples were taken as references. Titers were analyzed using the standard plaque assay. In this assay, Vero cells were infected with serial dilutions of viruses (10<sup>-1</sup> to 10<sup>-6</sup>) into duplicate wells.

After infection, the Vero monolayers were overlaid with 1X MEM (Sigma) supplemented with 2 mM L-Glutamine, 2.7% sodium bicarbonate, 5% HI FBS, 1% antibiotic/antimycotic solution [100 U/ml of penicillin, 0.1 mg/ml of streptomycin, 0.25 µg/ml Amphotericin (Sigma)], and 44% of 0.6% agarose (Sigma). Cells were incubated for 4 days at 37°C, 5% CO<sub>2</sub>, and were then overlaid with a second overlay containing 1X MEM supplemented with 2 mM L-Glutamine, 2.6% sodium bicarbonate, 2% HI FBS, 1% antibiotic/antimycotic solution, 44% of 0.6% agarose, and 3% of Neutral red solution (Sigma). The plaques were counted 24 hours after the addition of the second overlay to determine the titer of the virus defined in plaque forming unit (PFU) per milliliter.

Virus penetration assay according to Vlaycheva et al. (J. Virol. 76:6172-6184, 2002)

To demonstrate that the M-60 mutation (and E-107 mutation) facilitates penetration in SF Vero cells, SF Vero cells were infected with Clone A, C, and I viruses, appropriately diluted in SF medium, for 5, 10, 20, or 60 minutes, and then treated for 3 minutes with 0.1 M glycine, 0.1 M NaCl, pH 3.0, to inactivate extracellular virus. Wells were washed twice with PBS, and then monolayers were overlaid with methyl-cellulose, followed by staining plaques on day 5 with crystal violet. Efficiency of penetration was calculated as the percentage of observed plaque numbers after glycine treatment, as compared to control infected wells that were treated with PBS instead of glycine.

### Clinical trials of Chimeri $Vax^{TM}$ -JE

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A clinical study (protocol H-040-003) was performed. The vaccine

administered to healthy adult male and female subjects had the native sequence at

M60 (arginine). Healthy adult subjects/group received a subcutaneous dose of graded
doses of ChimeriVax<sup>TM</sup>-JE vaccine, and various control groups were included.

Eleven to 33 subjects were tested per dose group. Viremia was measured daily by
plaque assay in Vero cell monolayers. The same assay and laboratory determined
viremia levels in both trials.

Safety assessments included the recording of adverse events, body temperature, physical examination, and laboratory tests (including measurement of viremia levels). Viremia was seen in the majority of subjects receiving ChimeriVax<sup>TM</sup>-JE.

A second study (protocol H-040-007) was performed in healthy adult male and female subjects in which 31 or 32 subjects per group received graded subcutaneous doses (3, 4, or 5 log<sub>10</sub> PFU) of ChimeriVax<sup>TM</sup>-JE containing the M60 cysteine mutation. The dose range was similar to that in the previous study in subjects who had received 2.8, 3.8, and 4.8 log<sub>10</sub> PFU.

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#### Results

Adaptive mutations in uncloned SF ChimeriVax<sup>TM</sup>-JE virus, and preparation of cloned variants

A diagram of virus samples produced in this study is shown in Fig. 6. The initial uncloned passage 2 (P2) sample (Pre-Master Seed candidate; PMS) was obtained in SF culture by transfecting cells with *in vitro* RNA transcripts that had been used to produce the vaccine in FBS-containing media for previous studies (Monath et al., Vaccine 20:1004-1018, 2002) followed by an additional amplification passage. The full genome of this virus was sequenced and shown not to contain any detectable mutations (Table 7) (note that the consensus sequencing approach does not detect minor subpopulations; detection limit of mutations is ~ 10%). Small-scale passages starting from this P2 virus to P10 level were performed in T25 flasks to analyze its genetic stability (g.s.) during prolonged propagation in SF culture (Fig. 6; g.s. passages). The full genome sequences of the g.s. P5 and g.s. P10 passages had one nucleotide change from C to T at nucleotide 935 resulting in an R to C amino acid substitution at residue M-60 (Table 7). This mutation was first detectable as heterogeneity at the g.s. P4 passage, but not g.s. P3.

Despite the results of small-scale genetic stability analysis, when three large scale manufacturing SF passages were performed from the uncloned P2 PMS in roller bottles to produce candidate uncloned Master Seed (P3) and the Production Seed (P4), and then in 100 L bioreactors to produce vaccine bulk (P5), a different mutation accumulated, an F to L amino acid change at residue E-107 due to a T to C change at nucleotide 1301 observed as a 50:50% heterogeneity (Table 7). This was an

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unacceptable mutation because it is a reversion from the SA14-14-2 sequence to wild type JE sequence at a critical attenuating residue (Arroyo et al., J. Virol. 75:934-942, 2001) and thus could potentially compromise safety of the vaccine.

Based on considerations mentioned below, cloned PMS candidates were then generated by plaque purification, to stabilize the SF vaccine and prevent accumulation of undesirable mutations, such as E-107. Plaque purification removes random mutations in uncloned virus introduced by in vitro transcription characterized by low fidelity of RNA synthesis compared to viral RNA synthesis by YF 17D-specific RNA polymerase (Pugachev et al., J. Virol. 78:1032-1038, 2004). Starting from the uncloned P2 PMS virus, a biological clone at P7, Clone A virus, which did not have any amino acid substitutions was obtained by three sequential plaque purifications followed by two amplification passages in SF medium, and was designated nonmutant P7 Clone A PMS. Its genome contained two silent nucleotide changes, at nucleotides 6952 and 7147 (Table 7). These changes were acceptable because they did not change the amino acid sequence of viral proteins and were located outside cisacting RNA elements essential for efficient virus replication. A Clone C P10 virus containing the M-60 mutation (designated M-60 P10 Clone C PMS variant) was produced similarly starting from the P5 g.s. virus (Fig. 6). In addition to the desired M-60 mutation, it only contained a silent nucleotide change at nucleotide 3616 (Table 7). Additionally, research-grade Clone I and Clone E viruses were later also isolated from the uncloned P5 vaccine bulk virus by a single plaque purification (selecting large plaque) and one amplification passage in Vero cells. The Clone I contained a single amino acid change at the E-107 residue, which was a reversion to wild type from amino acid F to amino acid L. Thus, Clone I represents a pure population of the E-107 revertant. Clone E contained a single amino acid mutation at the N-terminus of the M protein, a Q to P amino acid change at residue M-5.

To ascertain genetic stability of the cloned PMS variants, relatively large scale g.s. passages mimicking manufacturing events were performed in SF culture (Fig. 6) (sequential passages designated S were done in T-225 flasks, and passages designates F were done in a 5 or 15 L bioreactor in which Vero cells were grown on Cytodex I microcarrier beads). Sequencing of the prM-E region only (cDNA Fragment I) was performed for the SSS and SSF samples (obtained by three Static passages, or two Static plus one Fermenter passages, respectively) of both candidates, and the FFF

sample of the M-60 variant. None of these g.s. samples had any detectable mutations in the prM or E proteins of the viruses other than the M-60 mutation in Clone C. There was no trace of the E-107 mutation (Table 7). This indicated that an acceptable level of genetic stability was achieved due to plaque-purification. The high genetic stability of the M-60 variant was subsequently confirmed during manufacturing of new Master (P11) and Production Virus (P12) Seeds produced in cell factories and final vaccine bulk (P13) produced in a 50 L bioreactor, all of which retained the M-60 mutation, but had no other detectable changes in their full genomes by consensus sequencing.

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Effects of the M-60 and E-107 mutations on virus growth in SF Vero cells

To compare growth kinetics of the non-mutant, M-60 mutant, and E-107 mutant viruses in SF culture, cells were infected at an MOI of 0.001 pfu/ml (confirmed by back-titration) with the uncloned P2 PMS, the uncloned P5 g.s. sample (M-60 mutant), or the uncloned P5 vaccine bulk variant (containing the E-107 mutation), as well as the uncloned P3 Master Seed and P4 Production Seed viruses also containing a proportion of the E-107 mutation. Daily aliquots of virus-containing media were harvested and titrated by plaque assay. As shown in Fig. 7, the M-60 virus grew faster than the non-mutant P2 virus and produced significantly (more than 10 times) higher titers on days 3 and 4 post-infection. The E-107 mutation also enhanced virus replication similarly to the M-60 mutation. Thus, both the M-60 and E-107 mutations clearly conferred a growth advantage in SF culture. In support of this conclusion, daily samples from the S, SSS, and SSF g.s. passages of both the nonmutant lone A and M-60 mutant clone C viruses (see Fig. 6) were collected and titered to analyze growth kinetics with the result that the M-60 mutant invariably produced up to 10 times higher peak titers (close to 8 log<sub>10</sub> pfu/ml) compared to nonmutant. Additionally, this conclusion was confirmed by comparing growth curves of Clones A, C, and I viruses in small scale SF culture, as Clones C (M-60) and I (E-107) invariably grew to higher titers than Clone A (non-mutant).

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Effects of the M-60 and E-107 mutations on neurovirulence of ChimeriVax<sup>TM</sup>-JE in suckling mice

Mouse neurovirulence tests have been used to ensure that neurovirulence of ChimeriVax<sup>TM</sup> vaccine candidates does not exceed that of the YF 17D vector. The YF 17D vaccine is lethal for mice of all ages after IC inoculation. In contrast, ChimeriVax<sup>TM</sup> vaccines are significantly more attenuated. Since adult mice generally are not sensitive to detect subtle differences in neurovirulence, e.g., those due to a single amino acid change, a more sensitive suckling mouse model using survival analysis can be used for that purpose (Guirakhoo et al., Virology 257:363-372, 1999; Guirakhoo et al., Virology 298:146-159, 2002; Monath et al., J. Virol. 76:1932-1943, 2002).

Eight day-old suckling mice were inoculated IC with serial dilutions of the clone A P7 virus, clone C P10 virus (M-60 mutation), uncloned P5 vaccine bulk (E-107 mutation), as well as a previously produced FBS-containing control ChimeriVax TM-JE virus (P5 Quality Control Reference Standard virus; no mutations), YF 17D positive control (YF-VAX®), or mock inoculated with diluent. Mortalities over a period of 21 days, median IC 50% lethal dose values (LD50), and average survival times (AST) of mice that died are shown in Table 8. As expected, YF-VAX® was highly neurovirulent. Inoculation of 2.4 log<sub>10</sub> PFU of this virus caused 100% mortality with a short AST of 8.8 days. Both the P7 non-mutant and P10 M-60 mutant clones were as highly attenuated as the original FBS-containing version of the chimera, with LD<sub>50</sub> values > 5 log<sub>10</sub> PFU and longer AST. Thus, the M-60 mutation does not change the highly attenuated phenotype of the vaccine in this animal model. The unclosed P5 vaccine bulk virus was significantly more virulent compared to the clones, with an IC LD<sub>50</sub> of 3.1 log<sub>10</sub> PFU, but was less virulent compared to YF-VAX<sup>®</sup>. Subsequently, manufacturing passages (Master Seed, Production Seed, and Vaccine bulk) of the cloned M-60 vaccine were examined in this test under GLP conditions, with similar results. This confirmed the high genetic/phenotypic stability that was achieved by plaque purification and the use of M-60 mutation.

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Analysis of safety and efficacy in nonhuman primates

### Experiment 1

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In this experiment, neurovirulence of Clone C (M-60 mutant)

ChimeriVax<sup>TM</sup>-JE Vaccine Master Viral Bank (MVB) and Production Viral Bank

(PVB) were compared after IC administration to cynomolgus monkeys, using YF
VAX® virus as a control (Table 9).

No vaccine-related clinical signs or changes in food consumption, body weight, or serum chemistry, and hematology parameters were observed. Lymphoid hyperplasia, consisting of increased size and number of lymphoid nodules in the spleen, was noted for 9 of 11, 4 of 11, and 8 of 11 monkeys from Groups 1-3, respectively. Although this finding is a common background finding in cynomolgus monkeys, the group incidences were greater than normal in these monkeys and were considered secondary to the expected immune response induced by the vaccines. It is noteworthy that similar changes occurred in both the ChimeriVax™-JE treatment groups and the YF-VAX® reference control group. [Some of the monkeys in all three groups developed low level postinoculation viremia of short duration, which was within acceptable limits, and all animals seroconverted to viruses used for inoculation. On Day 31, yellow fever virus-specific neutralizing antibody titers for the YF-VAX®treated monkeys ranged from 2.07 to >6.13 in the LNI assay, and no YF-VAX®treated monkeys had cross-reactive antibodies to JE virus in the PRNT50 assay. All ChimeriVaxTM-JE MVB vaccine-treated monkeys had JE neutralizing antibody titers ≥ 320 (range 320 to >20480) and had no cross-reacting antibody to YF virus in the LNI assay. All ChimeriVax™-JE PVB vaccine-treated monkeys had JE neutralizing antibody titers  $\geq$  160 (range 160 to >20480) and had no cross-reacting antibody to YF virus. There was no discernible relation between magnitude or duration of detectable viremia and the magnitude of JE-neutralizing antibody titer induction].

The ChimeriVax<sup>TM</sup>-JE MVB and PVB preparations exhibited minimal neurovirulence in this test. The most comprehensive measure of neurovirulence in the monkey neurovirulence test for Flavivirus vaccines is the combined group mean lesion score, representing the average of the mean target area and mean discriminator area scores. The target areas in cynomolgus monkeys are the substantia nigra and the cervical and lumbar enlargements of the spinal cord and represent regions of the central nervous system (CNS) that are injured by all Flaviviruses. The discriminator

areas are the globus pallidus, putamen, anterior and medial thalamic nuclei, and lateral thalamic nucleus, and represent regions of the CNS that are injured selectively by strains of YF 17D (and presumably other Flaviviruses) having different virulence properties, and that discriminate between a reference strain and a strain having increased neurovirulence. The combined mean lesion scores for monkeys treated with the ChimeriVax<sup>TM</sup>-JE MVB and PVB preparations were significantly lower than for the YF-VAX<sup>®</sup> reference control group (p<0.05). The mean discriminator center scores for the two groups of monkeys treated with the ChimeriVax<sup>TM</sup>-JE MVB and PVB were also significantly lower than for the YF-VAX<sup>®</sup> reference control group (p<0.05) (Table 9). There was no statistically significant difference between mean scores for the 2 groups of monkeys that received the ChimeriVax<sup>TM</sup>-JE vaccine preparations, and both preparations demonstrated similarly low neurovirulence in the monkey neurovirulence test.

Thus, the results of the monkey neurovirulence test show that the new (M60, Clone C) plaque-purified MVB and PVB have a satisfactory safety profile. The test articles displayed no clinical toxicity, and had significantly lower discriminator and combined lesion scores on neuropathological examination than the reference control (YF-VAX®). The test articles did not differ from the reference control (YF-VAX®) in viscerotropism, as measured by quantitative viremia.

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#### Experiment 2

This experiment was done to compare viremia, immune response, and safety of the original uncloned P5 ChimeriVax<sup>TM</sup>-JE Vaccine [produced previously in Vero cells in the presence of FBS, had no mutation except for E491 L to F change located in the hydrophobic tail of the E protein, which appears to be a benign mutation in terms of biological phenotype, and it has already been tested in clinical trials (Monath et al., J. Infect. Dis. 188:1213-1230, 2003; Monath et al., Vaccine 20:1004-1018, 2002)] and the new Clone C (M-60 mutant) ChimeriVax<sup>TM</sup>-JE purified vaccine bulk (P13) following a single subcutaneous (SC) administration in cynomolgus monkeys. ChimeriVax<sup>TM</sup>-JE virus was detected in the sera of 5 (100%) of 5 seronegative monkeys inoculated with original uncloned P5 ChimeriVax<sup>TM</sup>-JE vaccine. The

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duration of viremia was 2-5 days with titers ranging from 20 to 790 PFU/mL. The mean peak viremia (±SD) was 244 (±310) PFU/mL, and the mean number of viremic days was 3.4 (± 1.34) (Table 10).

ChimeriVax<sup>TM</sup>-JE virus was detected in the sera of 4 (100%) of 4 seronegative monkeys inoculated with the new P13 JE vaccine purified bulk. The duration of viremia was 2-5 days with titers ranging from 50 to 290 PFU/mL. The mean peak viremia (±SD) was 160 (±123) PFU/mL, and the mean number of viremic days was 3.75 (±1.26) (Table 10). Neither mean peak viremia nor number of viremic days differed significantly between the two treatment groups (p-values 0.6290 and 0.7016, respectively; ANOVA). 10

All seronegative monkeys seroconverted following treatment with the original uncloned P5 ChimeriVax™-JE Vaccine or P13 JE Vaccine Purified Bulk (Table 10). On Day 31, sera from 5 (100%) of 5 monkeys inoculated with uncloned P5 Vaccine had JE virus neutralizing antibody titers ranging from 640 to 5120 (geometric mean titer = 1689). Sera from 4 (100%) of 4 monkeys inoculated with P13 ChimeriVax<sup>TM</sup>-JE Vaccine Purified Bulk had JE virus neutralizing antibody titers ranging from 320 to 2560 (geometric mean titer = 761). Antibody titers did not differ significantly between treatment groups (p = 0.2986, ANOVA).

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Thus, the new M-60 vaccine was compared to the original uncloned ChimeriVax™-JE vaccine (no mutations except for E491) with respect to safety (viremia) and immunogenicity. The new vaccine was slightly less viscerotropic (a desirable feature) but still highly immunogenic. The differences in the magnitude of viremia and immunogenicity were not statistically significant.

Effects of M-5, M-60, and E-107 mutations on the pH threshold of virus infectivity ChimeriVaxTM-JE vaccine was produced by insertion of prM and E genes from SA14-14-2 strain of JE virus into backbone of YF 17D virus. The envelope of SA14-14-2 virus (present in ChimeriVaxTM-JE) differed from its parent SA14 virus by 10 amino acids: E107 L to F, E138 E to K, E176 I to V, E177 T to A, E227 P to S, E244 E to G, E264 Q to H, E279 K to M, E315 A to V, and E439 K to R (Guirakhoo 30 et al., Virology 257:363-372, 1999). By site-directed mutagenesis it was shown that some of these residues were involved in attenuation of ChimeriVax<sup>TM</sup>-JE virus.

Mutants or revertants of ChimeriVax<sup>TM</sup>-JE were selected to identify whether mutations have altered the pH threshold of these viruses. To determine whether the M-60, E-107, or M-5 mutations affect virus infectivity in a pH-dependent fashion, a standard assay for pH threshold of infectivity was performed as described in Materials and Methods. The following viruses were tested: (1) ChimeriVax<sup>TM</sup>-JE non-mutant (clone A, P7 containing all 10 SA14-14-2 mutations in the E protein); (2) ChimeriVax<sup>TM</sup>-JE E107 F to L revertant (clone I P6, containing 9 E protein mutations); (3) ChimeriVax<sup>TM</sup>-JE M60 R to C mutant (clone C, P10 containing all 10 E protein mutations), and (4) M-5 Q to P mutant (clone E, P6 containing all 10 E protein mutations) (Table 12).

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Non-mutant clone A P7 virus, M-60 mutant clone C P10 virus, M-5 mutant clone E, and uncloned P5 virus containing the E-107 mutation were treated with a range of decreasing pHs followed by titration of residual viral infectivity. Infectivity of three viruses (clone A control virus, Clone C M60 mutant, and Clone I E-107 mutant) started to drop uniformly after pH 6.0 and was lost at pH 5.8 (pH threshold 5.9), except for M5 mutant Clone E virus. The M-5 mutant had a significantly higher pH threshold (pH 6.3) compared to all other viruses (pH 5.9) (Fig. 8A). This is the first direct evidence that the ectodomain of M protein plays an essential role in the process of infection of cells by a Flavivirus. Thus, the N-terminus of M protein may function in the process of fusion triggered by a low pH in endosomes following virus adsorption and internalization, which is a function attributed previously solely to the envelope E protein.

The pH threshold of 5.9 for fusion of ChimeriVax<sup>TM</sup>-JE viruses is lower than those described for other wild-type (wt) Flaviviruses (Guirakhoo et al., J. Gen. Virol. 72:1323-1329, 1991) and may be involved in attenuation of the virus.

These data demonstrated that the E-107 mutation in the E region of ChimeriVax<sup>TM</sup>-JE did not change the pH threshold for fusion. Generally, a low pH threshold means that more protonization of specific amino acids is required for conformational changes in the E-protein to occur that are necessary for transition from dimer to trimer. It is likely that one or more SA14-14-2 specific mutations (other than the E107 mutation, which is located within the conserved fusion peptide) are responsible for retaining the low pH threshold (pH 5.9) for fusion and consequently attenuated phenotype of the virus for the host. Apparently, the M-5 mutation is

capable of increasing this threshold from 5.9 to 6.3, which is closer to those of wt
Flaviviruses (Guirakhoo et al., Virology:169(1):90-99, 1989; Guirakhoo et al., J. Gen.
Virol. 72:1323-1329, 1991). An increase in pH threshold for fusion should
theoretically decrease the attenuated phenotype of the virus, because the viruses can

fuse at higher pHs with less protonization required for transition to a fusion active
state. This appeared to be true, since M5 virus inoculated at 1.4 log<sub>10</sub> PFU into 3-4
day old suckling mice by the intracerebral route was significantly more virulent than
the control virus (ChimeriVax<sup>TM</sup>-JE vaccine virus without the M5 mutation)
inoculated at 1.7 log<sub>10</sub> PFU (p=0056) (Fig. 8B). Nevertheless, the M5 mutant virus

(at a dose of 1.4 log<sub>10</sub> PFU) remained significantly less neurovirulent than YF-VAX<sup>®</sup>
(at a dose of 0.9 log<sub>10</sub> PFU) in 3-4 day old suckling mice (Fig. 8C), indicating than the
SA14-14-2 mutations within the envelope protein of the vaccine virus are still
providing a sufficient level of attenuation for this virus.

### 15 Mutations in other chimeras that affect pH threshold for fusion

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The Indirect Fusion Assay was performed using two groups of each ChimeriVax<sup>TM</sup>-DEN vaccines viruses: ChimeriVax<sup>TM</sup>-DEN1-4 P7 containing no É protein mutations and ChimeriVax<sup>TM</sup>-DEN1-4 P10 which contained single mutations in the E protein, except for ChimeriVax<sup>TM</sup>-DEN4 P10. Viruses were incubated with media of different pH for 10 minutes at room temperature. The titers were determined, after returning the pH to the neutral pH, using a standard plaque assay. As shown in Table 13, the threshold for virus inactivation (fusion) was similar between P7 and P10 of ChimeriVax<sup>TM</sup>-DEN2 and DEN4 viruses (pH 6.4). In contrast, the pH threshold for ChimeriVax<sup>TM</sup>-DEN1 P10 was 0.4 units lower than that of ChimeriVax<sup>TM</sup>-DEN1 P7 virus (pH 6.0 vs. pH 6.4). The difference in pH threshold was less dramatic for ChimeriVax<sup>TM</sup>-DEN3 P10 virus (pH 6.4 vs. pH 6.2).

The maximum virus inactivation occurred at pH 6.2 for all P7 of ChimeriVax<sup>TM</sup>-DEN viruses except for that of ChimeriVax<sup>TM</sup>-DEN4, which was slightly lower (pH 6.0). It appeared that ChimeriVax<sup>TM</sup>-DEN1 P10 required a significantly lower pH for complete inactivation (pH 5.6). Both ChimeriVax<sup>TM</sup>-DEN1 and -DEN3 viruses contain an amino acid substitution at E-204 from K to R (the E-protein of DEN3 is 2 amino acids shorter than other 3 serotypes, therefore, the E-202 residue in this virus is homologous to E-204 in DEN1). The less dramatic

difference in fusion threshold for the DEN3 chimera might be due to presence of WT (K) and mutant R amino acids (E204K/R) in P10 virus stock as was shown by consensus sequencing (K:R=50:50) (Pugachev et al., J. Virol. 78:1032-1038, 2004). Since no change in threshold for virus inactivation was observed with DEN2 P10 chimera, despite the E251 mutation, it can be concluded that the mutation at this residue is not involved in viral fusion process (Fig. 8D).

In order to determine if the presence of K/R heterogeneity in P10 of ChimeriVax<sup>TM</sup>-DEN3 was responsible for its non-dramatic change in pH threshold for fusion, the indirect fusion assay was performed using P7 (no mutation, E202K), P10 (50% mutation, E202K/R), and P15 (complete mutation, E202R) viruses. As shown in Fig. 8E, the pH threshold for inactivation (fusion) of ChimeriVax<sup>TM</sup>-DEN3 P10 was at pH 6.2, which was between those for ChimeriVax<sup>TM</sup>-DEN3 P7 (pH 6.4) and ChimeriVax<sup>TM</sup>-DEN3 P15 (pH 6.0) viruses. Since the E202 K to R mutation was the only amino acid substitution detected in E-protein of these chimeras, it is most likely that this mutation is responsible for a 0.4 pH shift in pH threshold for fusion of the P15 virus.

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As mentioned above, the E204 K to R mutation, which occurred during cell culture manufacture of the vaccine, lowered the pH threshold for fusion by 0.4 units of pH. The E204 K to R mutation appears to generate new intramolecular H bonds and a new salt bridge, which might have a significant impact on the dissociation of the E dimers. The structure of the ChimeriVax<sup>TM</sup>-DEN1 (PMS, P7) E protein was modelled based on the atomic coordinates of 394 residues of the DEN2 E-protein ectodomain (S1 strain) determined in the presence of the detergent n-octyl-β-Dglucoside (Modis et al., Proc. Natl. Acad. Sci. U.S.A. 100:6986-6991, 2003). The K residue at position 204 was changed to R to mimic the mutant virus, and the modelling was repeated to represent the E-protein structure of the ChimeriVaxTM-DEN1 (VL, P10) virus (Guirakhoo et al., J. Virol. 78:9998-10008, 2004). The K residue at position 204 (204K) is located within a short loop, in a hydrophobic pocket lined by residues, which have been shown to influence neurovirulence or the pH threshold for fusion (Lee et al., Virology 232:281-290, 1997; Lindenbach et al., 2001 Flaviviridae: the viruses and their replication. Fields Virology, eds. Knipe D.M., and Howley P.M. [Lippincott Williams and Wilkins, Philadelphia], 1, 991-1004; Monath et al., J. Virol. 76:1932-1943, 2002). In Fig. 8F, the homology model of the E-

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homodimer structure of the vaccine virus (204R) is compared to that of the PMS (204K) virus. The side chains of 204K and 261H of one of E monomer appeared to make H bonds with the backbone atoms of 252V and 253L residues, respectively, on the opposite monomer. At position 204, the R in the E protein of the vaccine virus (VL P10) is predicted to reorient itself so that these hydrogen (H) bonds are lost. Instead the side chain of the mutant R is in proximity with 261H and 257E, resulting in the generation of new intramolecular H bonds between 204R and 261H, and probably of a new salt bridge between 204R and 257E. Since the pk of Histidine could be approximately 6.0, which is slightly below the fusion threshold (pH ~6.4), the initial hypothesis by Guirakhoo et al., (J. Virol. 78:9998-10008, 2004) was that the predicted new H bonds between 204R and 261H and the salt bridge between 204R and 257E, might affect the pH threshold of fusion. This theory turned out to be true, since the experiments described here revealed that the threshold for fusion of ChimeriVax<sup>TM</sup>-DEN1 is around 6.0, which is 0.4 pH units lower than its P7 virus (pH 6.4). Apparently, the new intermolecular bonds introduced by R at residue 204 strengthen the association of the E-dimer so that the transition to low pH requires more protonization of appropriate residues (e.g., H 261). The lower threshold for fusion affects viscerotropism of the virus in monkeys and reduces neurovirulence for suckling mice inoculated by the i.c. route (Guirakhoo et al., J. Virol. 78:9998-10008, 2004).

The E202 K to R substitution in the E-protein of the ChimeriVax<sup>TM</sup>-DEN3 P10 vaccine is homologous to the E204 mutation in the ChimeriVax<sup>TM</sup>-DEN1 P10 vaccine. As with ChimeriVax<sup>TM</sup>-DEN1 P10, ChimeriVax<sup>TM</sup>-DEN3 P10 (heterogenous at residue 202 containing both K and R residue) showed a lower pH threshold (~0.2 pH unit) for fusion when compared to P7. The pH threshold for fusion was further lowered (0.4 pH unit, similar to ChimeriVax<sup>TM</sup>-DEN1 P10) when the mutation was fixed at P15 of ChimeriVax<sup>TM</sup>-DEN3. This data showed that the residue 202/204 may be a universal determinant of attenuation in all dengue viruses. Currently, ChimeriVax<sup>TM</sup>-DEN3 and -DEN4 P10 vaccine viruses do not contain this mutation and both viruses induce a higher viremia levels in monkeys (Guirakhoo et al., J. Virol. 78:4761-4775, 2004) inoculated with a tetravalent vaccine formulation. It remains to be seen if K to R mutation in ChimeriVax<sup>TM</sup>-DEN3 or ChimeriVax<sup>TM</sup>-DEN4 would lower their viscerotropism in their hosts.

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It was previously reported that WT-JE had a pH threshold for fusion of 6.4 (Guirakhoo et al., J. Gen. Virol. 72:1323-1329, 1991). In this study, all variants of ChimeriVax<sup>TM</sup>-JE had a pH threshold of 5.9. The low pH threshold observed in these experiments is likely due to the presence of one or more of the 10 attenuating mutations in the envelope protein of ChimeriVax TM-JE. This mutation might strengthen the association of the E-protein dimer so that a lower pH is required for dissociation and transition to trimer structure and subsequent fusion. The data presented here showed that neither the E107 F to L mutation (located in the cd-loop of the domain II of the E-protein) nor the E279 M to K mutation (located within the hydrophobic pocket of the domain II) was responsible for lowering the pH threshold. It is possible that other mutations in the JE E protein may affect the pH threshold for fusion. Analysis of the crystal structure of TBE virus E protein, which closely resembles the JE E protein, can help to predict the residues that, if altered, could change the pH threshold for fusion. Based on this model, it is likely that the mutations in residues E244 G and/or E264 H are responsible for a lower pH threshold, than the WT JE, for fusion of ChimeriVax<sup>TM</sup>-JE virus.

Effect of the M-60 and E-107 mutations on efficiency of virus penetration

The effects of the M-60 (Clone C virus) and E-107 (Clone I virus) mutations on virus penetration into SF Vero cells were examined using the method of Chambers (Vlaycheva et al., J. Virol. 76:6172-6184, 2002). In this experiment, SF Vero cells were infected with appropriately diluted viruses (to yield ~ 50 plaques/well at each time point) for 5, 10, 20, or 60 minutes. Un-internalized virus is inactivated by addition of acidic glycine silution, while control parallel wells are treated with PBS (neutral pH). Cells are washed with PBS and overlaid with methyl-cellulose overlay, followed by visualization and counting of plaques on day 5. The efficiency of penetration is presented as a percentage of the average number of plaques in glycine-treated wells relative to the number of plaques in control, PBS treated wells. A preliminary penetration test result is shown in Fig. 9A. It is important that the percentages of penetrated Clone C and Clone I viruses were higher than the non-mutant Clone A virus at 5 and 10 minute time points, at which effects of mutations on penetration are more likely to be detected. The result is not statistically significant as evidenced by standard deviation bars and needs to be confirmed in additional repeat

tests. Nevertheless, this experiment suggested that both the M-60 and E-107 mutations could improve the efficiency of membrane fusion of ChimeriVax<sup>TM</sup>-JE virus to cells grown in SF conditions. A possible mechanism of the effect of the M-60 and E-107 residues on process of membrane fusion is illustrated in Fig. 9B. The M-60 residue is located in the viral membrane, while the E-107 residue inserts into the cell membrane, and the two membranes are forced to fuse following low pH-dependent rearrangement of the E protein (which based on our data could be facilitated by the M protein ectodomain). A more appropriate amino acid at either of these two residues may facilitate fusion of the membranes.

Because our data establish for the first time that both the ectodomain of the M protein and its transmembrane domain are of functional significance, the entire M protein can now be considered an attractive target for mutagenesis to attenuate Flaviviruses for the purpose of developing new live attenuated vaccines. For example, random or specific (following further analysis of protein structure) amino acid changes, or deletions of increasing length, e.g., of 1, 2, 3, 4, 5, etc., amino acids, can be incorporated throughout the protein with the expectation that biological phenotype of the virus will be altered, resulting in significant attenuation.

#### Results from clinical trial

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The viremia profiles of ChimeriVax<sup>TM</sup>-JE with the arginine and cysteine M60 residues as obtained from the clinical trials noted above are compared in Tables 11 A and B. In subjects receiving ChimeriVax<sup>TM</sup>-JE M60 arginine, 67-100% of the subjects were viremic on one or more days, compared to 29-50% for subjects receiving ChimeriVax<sup>TM</sup>-JE M60 cysteine. The mean maximum viremia levels in subjects receiving ChimeriVax<sup>TM</sup>-JE M60 arginine ranged from 13 to 40 PFU/ml, compared to mean maximum viremia levels of 3.5-6.3 PFU/ml in the case of ChimeriVax<sup>TM</sup>-JE M60 cysteine. The duration of viremia was also notably longer in the case of ChimeriVax<sup>TM</sup>-JE M60 arginine.

These data demonstrated that the level of viremia is notably lower in the case of the vaccine containing the M60 mutation. Viremia is a measure of viscerotropism (virulence) of the vaccine virus. A vaccine with reduced viremia is considered safer, since cell damage and dysfunction of organs sustaining virus replication and contributing to viremia is reduced, as is the likelihood that the virus will cross the

blood brain barrier and invade the central nervous system. In other experiments, it was shown that the M60 mutant was as highly immunogenic in humans as the non-mutant.

Table 1. Consensus sequence of small plaque (P10 PMS) (P/N IT-0116; L/N I020504A) (plaque purified from p5 Run 1 Vaccine Lot).

Position	Amino Acid change	NT position	NT change
M(66)	Leucine → Proline	954	CTA → CCA
E(313)	Glycine → aRginine	1919	GGG → AGG
	Asparagine (silent)	2926	AAC → AAT
	Glycine (silent)	7126	GGA → GGG

Table 2. Consensus sequence of large plaque PMS (P10, PMS) (P/N IT-0117; L/N 1030804A) (derived from p5 Run 1 Vaccine Lot).

Position	Amino Acid change	NT position	NT change
E(313)	Glycine → aRginine	1919	GGG → AGG
	Glycine (Silent)	7126	GGA → GGG

Table 3. Sequence of large plaques isolated after 2 additional passages of the S plaque PMS (p10) in Vero cells under serum free conditions.

LP Isolate	Position	Amino Acid Change	NT #	NT change	Immuno- Stain
110, 1117	3466	Leucine → Proline	954	CTA →	SP
#3, #7,	M66	Leucine - Fromie		CCA	
<u>#8, #9,</u>		ļ			
#10, #11,				•	
#12, #13,					
#14, #18,					ļ
#19, #20					
#1	M62	Valine → Methionine	941	TGT → TAT	SP
	M66	Leucine → Proline	954	CTA → CCA	
#2	M62	Valine → Glycine	942	GTG → GGG	SP
		Valine → Glutamic Acid	942	GTG → GAG	]
	M66	Leucine → Proline	954	CTA → CCA	
#4	M63	Phenylalanine → Serine	945	$TTT \rightarrow TCT$	LP
#5	M62	Valine → Alanine	942	GTG →	SP
	M66	Leucine → Proline		GCG	-
#6	M66	Leucine → Proline	954	CTA → CCA	SP
	M64	Valine (Silent)	949	GTC → GTT	
#15	M62	Valine → Alanine	942	GTG → GCG	SP
	M66	Leucine → Proline	954	CTA → CCA	
#16	wt	Leucine	N/A	CTA	LP/SP
"2"	M66	Leucine → Proline	954	CTA → CCA	
#17	M64	Valine → Isoleucine	947	GTC → ATC	SP
	M66	Leucine → Proline	954	CTA → CCA	-

Table 4. Observed CPE for HepG2.

			!						
Days Post	0	1	2	3	4	5	6	7	8
Infection									
WN01	0%	0%	0%	0%	0%	30%_	90%	~100%	100%
WN02 P5	0%	0%	0%	5%	30%	50%	~100%	100%	
WNL	0%	0%	0%	0%	0%	30%	90%	~100%	100%
WNS	0%	0%	0%	5%	30%	50%	~100%	100%	
YF/17D	0%	0%	0%	20%	50%	70%	~100%	100%	
IF/I/D	0/0	1 0/0	1 0 / 0	1-3/0	1 2 3 7 0				

### Table 5. Viremia in monkeys inoculated with ChimeriVax $^{TM}$ -WN02 vaccine or YF-VAX $^{\otimes}$ . 5

Treatment Group	Monkey Number	Day 1**	Day 2	Day 3	Day 4
			ŀ	·	
YF-Vax®	MF21157M	0	0	20	200
YF-Vax®	MF21214F	0	0	0	0
YF-Vax®	MF21151M	- 0	. 0 -	10	60
YF-Vax <sup>®</sup> YF-Vax <sup>®</sup>	MF21252F	0	0	0	0
ChimeriVax TM-WN					000
Vaccine (P5)	MF2808M	0	30	790	820
ChimeriVax <sup>1M</sup> -WN Vaccine (P5)	MF21205F	0	50	160	100
ChimeriVax <sup>TM</sup> -WN Vaccine (P5)	MF21139M	0	10	180	70
ChimeriVax M-WN Vaccine (P5)	MF21191F	0	80	970	1000

<sup>\*</sup>Viremia expressed as pfu/mL

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<sup>\*\*</sup>Day 1: Study Day 1, monkeys inoculated on Study Day 1
Zero PFU/mL means below the limit of detection, theoretical assay cutoff = 10 PFU/mL

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Table 6. Sequence of the M region of YF-WN chimera obtained directly from a plaque isolate from viremic monkeys inoculated with WN02 vaccine virus.

Monkey #	Day of viremia	Plaque Isolate #	Visible Plaque Morphology (at time of picking)	M66 Present?	Additional M Mutations
21205	4	#4	SP	NO	NO
2808	1 3	#8	SP	NO	NO
2808	3	#9	LP	NO	M60 (R to G)
21191	2	#10	LP	NO	NO
	1	#14	SP	NO	M61 (V to A)
21191	1 1	#15	SP	NO	NO
21191	1 1	#16	LP	NO	M63 (F to S)
21191	<u> </u>	#10	1-1		

Table 7. Nucleotide and amino acid sequences of the uncloned and cloned SF ChimeriVax<sup>TM</sup>-JE samples (see Fig. 6).

Candidate	Passage	Part of genome sequenced	Protein - a.a. No.b	Nt No.ª	Nucleotide change/ heterogeneity	Amino acid change/ heterogen.	Comments
Uncloned	P2 (PMS)	Full genome		-	-		No mutations
	P3 g. s. from PMS	a.a. M-60 only	•	-	-	-	No M-60 mutation  M-60 mutation first
	P4 g. s. from PMS	a.a. M-60 only	M-60	935	c/ T	R/C	detectable and dominant
	P5 g. s. from PMS	Full genome	M-60	935	C to T	R to C	M-60 mutation located in the cytoplasmic hydrophilic stretch of the M protein
	P10 g. s. from PMS	95% full genome	M-60	935	C to T	R to C	M-60 is the only detected rnutation
	cGMP P3 (MS)	prM-E	E-107	1301	T/c	F/L	Reversion to WT first detectable
	eGMP P4 (PS) Baxter	prM-E	E-107	1301	T/c	F/L	Reversion to WT
	cGMP P5 (VB) Baxter	Full genome	E-107	1301	T/C	F/L	Reversion to WT (~50%).
M-60 mutant	P10	Full	M-60	935	C to T	R to C	Desired/expected
clone C	PMS	genome	NS2A-26	3616	A to G		Silent No subpopulations detected
	SSS P13 g.s.	prM-E	M-60	935	C to T	R to C	
	SSF P13	prM-E	M-60	935	CtoT	R to C	No subpopulations detected
	g.s. FFF P13	ргМ-Е	M-60	935	C to T	R to C	No subpopulations detected
	g.s.	Full	NS4B-12	6952	C to T	<del></del>	Silent
Non-mutant	P7 PMS	genome	NS4B-77	7147	T to C		Silent
clone A	SSS P10	prM-E				•	No subpopulations detect
	g.s. SSF P10 g.s.	prM-E	-		-	T	No subpopulations detect

<sup>&</sup>lt;sup>a</sup>: From the beginning of the genome <sup>b</sup>: From the N-terminus of indicated protein

Table 8. Neurovirulence of clone A P7, clone C P10, uncloned P5, FBS-containing standard, and YF-VAX® viruses in 8 day-old suckling mice.

Virus	a.a. change	Dilution	Inoculation Dose Log <sub>10</sub> PFU	Mortality No. dead/No. inoculated (% mortality)	LD <sub>50</sub> Log <sub>10</sub> PFU	AST days
Clone A P7 PMS	None .	Neat _ 10 <sup>-1</sup> 10 <sup>-2</sup> 10 <sup>-3</sup> 10 <sup>-4</sup>	5.1 4.1 3.1 2.1	1/11 (9%) 3/11 (27%) 1/10 (10%) 1/12 (8.3%) 0/12 (0%)	>5.1	11 14 14 11 N/A
Clone C P10 PMS	M-60	Neat 10 <sup>-1</sup> 10 <sup>-2</sup> 10 <sup>-3</sup>	5.5 4.5 3.5 2.5 1.5	2/11 (18%) 0/10 (0%) 1/12 (8.3%) 0/12 (0%) 0/12 (0%)	>5.5	11 N/A 13 N/A N/A
Uncloned P5 VB	E-107	Neat 10 <sup>-1</sup> 10 <sup>-2</sup> 10 <sup>-3</sup>	5.3 4.3 3.3 2.3 1.3	9/10 (90%) 10/11 (91%) 9/11 (82%) 1/11 (9%) 1/10 (10%)	3.1	9.4 10.7 11.8 14
FBS- containing standard virus	none	Neat 10 <sup>-1</sup> 10 <sup>-2</sup> 10 <sup>-3</sup>	5.3 4.3 3.3 2.3	0/10 (0%) 0/10 (0%) 2/9 (22%) 0/11 (0%)	>5.3	N/A N/A 16.5 N/A
YF-VAX®	N/A	10-1	2.4	10/10 (100%)	<2.4	8.8
Sham (MEM- 10%FBS)	N/A	N/A	N/A 	0/10 (0%)	N/A	N/A

Table 9. Neurovirulence for Cynomolgus Monkeys of M-60 (Clone C) Master and Production seeds vs. YF-VAX $^{\otimes}$  control.

		Treatment.	Dose	E ;	sion scores (gro	oup mean; SD (ran
No:	*Male/Fe *male**		(PFU /0:25 ml/ inoculum)	Target	Discriminat or Areas	Combined
1	6/5	YF-VAX® (Commercial Yellow Fever Vaccine)	5.5 x 10 <sup>4</sup>	0.436 SD 0.190 (0.25-0.81)	0.610 SD 0.417 (0.25-1.38)	0.526 SD 0.194 (0.29-0.87)
2	5/6	ChimeriVax™-JE Vaccine Master Viral Bank P11 (M-60)	1.0 x 10 <sup>5</sup>	0.196 SD 0.210 (0-0.56)	0.183 SD 0.177 (0-0.44)	0.191 SD 0.163 (0-0.47)
3	6/5	ChimeriVax <sup>TM</sup> -JE Vaccine Production Viral Bank P12 (M-60)	1.0 x 10 <sup>5</sup>	0.223 SD 0.349 (0-0.56)	0.106 SD 0.138 (0-0.31)	0.167 SD 0.231 (0-0.63)

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PFU = plaque-forming units
24 of 11, 2 of 11, and 1 of 11 animals in groups 1, 2, and 3, respectively, were excluded from score calculations because they were found to be JE-seropositive on day I (pre-inoculation) in a retrospective PRNT50 test, which is more sensitive than HAI test used for prescreening.

Table 10. Comparison of magnitudes of viremia and immunogenicity in cynomolgus monkeys inoculated SC with the original uncloned P5 ChimeriVax<sup>TM</sup>-JE vaccine produced in FBS-containing medium (containing no mutations except for E491) and the new Clone C P13 purified vaccine bulk (M-60 mutant).

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	Sample		Sample Dose (PFU)		Viremia <sup>1</sup>		
Group No.	Number of Male/Fe- male			Mean peak titer ± SD (PFU/ml)	Mean duration ± SD (days)	on day 31 (geometric mean PRNT <sub>50</sub> titer (min., max)) <sup>1</sup>	
1	3/3	Diluent	0	0	0	N/D	
2	3/3	ChimeriVax <sup>TM</sup> -JE original uncloned P5 Vaccine	1.0 x 10 <sup>4</sup>	244 ± 310	3.4 ± 1.34	1689 (640, 5120	
3	3/3	Clone C (M-60) ChimeriVax <sup>TM</sup> -JE vaccine, purified bulk, P13	1.0 x 10 <sup>4</sup>	160 ± 123	3.75 ± 1.26	761 (320, 2560)	

<sup>&</sup>lt;sup>1</sup>2 of 6, 1 of 6, and 2 of 6 animals in groups 1, 2, and 3, respectively, were excluded from calculations of the values because they were found to be JE-seropositive on day 1 (pre-inoculation) in a retrospective PRNT50 test, which is more sensitive than HAI test used for prescreening.

Table 11A. Viremia profiles in subjects enrolled in Study H-040-003 in which ChimeriVax<sup>TM</sup>-JE with the M60 arginine amino acid was administered. The dose range in bold is similar to that given in another study (H-040-007) in which the mutant M-60 cysteine vaccine was administered.

Viremia		oseLognPBU(	himeriVax e	ue M60)arginin	
	518	4.8	3:8	2.8	1.8
	(0,10)	(n=33)	(n=1ii)		(0-111)
Viremic on 1 or more	5/10	22/33	9/11	11/11	9/11
days [No. viremic/total (%)]	(50%)	(67%)	(82%)	(100%)	(82%)
Mean peak viremia (PFU/mL)	7.0	13.0	16.4	40.9	18.2
Range in peak viremia (PFU/mL)	0 - 20	0 - 40	0 - 50	0 - 220	0 - 50
Mean duration (days)	0.9	1.6	1.4	2.7	2.2
Range in duration (days)	0 - 4	0 - 5	0 - 3	1 - 6	0 - 5

Table 11B. Viremia profiles in subjects enrolled in Study H-040-007 in which ChimeriVax<sup>TM</sup>-JE with the M60 cysteine amino acid was administered.

Viremia	Dose Logio	FU Glimenya cysteine (± 410 N-32	3:0 3:0 N 32
Viremic on 1 or more	9/31	16/32	13/32
days [No. viremic/total (%)]	(29%)	(50%)	41%)
Mean peak viremia (PFU/mL)	3.5	6.3	4.4
Range in peak viremia (PFU/mL)	0-20	0-30	0-10
Mean duration (days)	0.3	0.8	0.6
Range in duration (days)	0-2	0-4	0-3

Table 12. Values of pH threshold for fusion found with the fusion assay for each ChimeriVax<sup>TM</sup>-JE vaccine.

Virus 200	pH threshold for fusion
ChimeriVax TM-JE parent, clone A P7 (contains all 10 E mutations)	5.9
ChimeriVax TM-JE clone C P10 (M60 R to C mutant, contains all 10 E mutations)	5.9
ChimeriVax <sup>TM</sup> -JE clone I P6 (E107 F to L revertant, contains 9 E mutations)	5.9
ChimeriVax TM-JE clone E P6 (M5 Q to P mutant, contains all 10 E mutations)	6.3

# Table 13. Values of pH threshold for fusion found with the indirect fusion assay for each couple of ChimeriVax<sup>TM</sup>-DEN P7 and P10.

Virus	pH Threshold for fusion
ChimeriVax™-DEN1 PMS P7	6.4
ChimeriVax™-DEN1 VL P10	6.0
ChimeriVax™-DEN2 PMS P7	6.4
ChimeriVax™-DEN2 VL P10	6.4
ChimeriVax™-DEN3 PMS P7	6.4
ChimeriVax™-DEN3 VL P10	6.2
ChimeriVax™-DEN4 PMS P7	6.4
ChimeriVax™-DEN4 VL P10	6.4

Table 14
Engineering of YF/Flavivirus chimeras

5	Virus	Chimeric C/prM junction <sup>1</sup>	Chimeric E/NS1 junction <sup>2</sup>	5' ligation <sup>3</sup>	3' ligation <sup>4</sup>	Sites <sup>5</sup> eliminated or (created)
	YF/WN	X-cactgggagagcttgaaggtc (SEQ ID NO:1)	aaagccagttgcagccgcggtttaa (SEQ ID NO:2)	Aatll	Nsil	٠
10	YF/DEN-I	X-aaggtagactggtgggctccc (SEQ ID NO:3)	gatecteagtaceaacegeggtttaa (SEQ ID NO:4)	Aatii	Sphi	Sph1 in DEN
	YF/DEN-2	X-aaggtagattggtgtgcattg (SEQ ID NO:5)	aacceteagtaccaccegeggtttaa (SEQ ID NO:6)	Aatli	SphI	
15	YF/DEN-3	X-aaggtgaattgaagtgctcta (SEQ ID NO:7)	acccccagcaccacccgcggtttaa (SEQ ID NO:8)	Aatli	SphI	XhoI in DEN (SphI in
	DEN) YF/DEN-4	X-aaaaggaacagttgttctcta	accegaagtgtcaaccgcggtttaa (SEQ ID NO:10)	Aatli	Nsil	
20	YF/SLE	(SEQ ID NO:9)  X-aacgtgaatagttggatagtc (SEQ ID NO:11)	accettegtcgcacccgcggtttaa (SEQ ID NO:12)	Aatll	Sphl	Aatll in SLE
	YF/MVE	X-aatttcgaaaggtggaaggtc (SEQ ID NO:13)	gaccggtgtttacagccgcggtttaa (SEQ ID NO:14)	a Aatli	Agel	(Agel in YF)
	YF/TBE	X-tactgcgaacgacgttgccac (SEQ ID NO:15)	actgggaacctcacccgcggtttaa (SEQ ID NO:16)	Aatii	Agel	(Agel in YF)

1,2: The column illustrates the oligonucleotide used to generate chimeric YF/Flavivirus primers corresponding to the C/prM or E/NS1 junction. (See text). X = carboxyl terminal coding sequence of the YF capsid. The underlined region corresponds to the targeted heterologous sequence immediately upstream of the NarI site (antisense - ccgcgg). This site allows insertion of PCR products into the Yfm5.2 (NarI) plasmid required for generating full-length cDNA templates. Other nucleotides are specific to the heterologous virus. Oligonucleotide primers are listed 5' to 3'.

<sup>3,4:</sup> The unique restriction sites used for creating restriction fragments that can be isolated and ligated *in vitro* to produce full-length chimeric cDNA templates are listed. Because some sequences do not contain convenient sites, engineering of appropriate sites is required in some cases (footnote 5).

<sup>5:</sup> In parentheses are the restriction enzyme sites that must be created either in the YF backbone or the heterologous virus to allow efficient in vitro ligation. Sites not in parentheses must be eliminated. All such modifications are done by silent mutagenesis of the cDNA for the respective clone. Blank spaces indicate that no modification of the cDNA clones is required.

NGTAAATCCT GTGTGCTAAT TGAGGTGCAT TGGTCTGCAA ATCGAGTTGC TAGGCAATAA ACACATTTGG ATTAATTTTA 41 ATCGTTCGTT GAGCGATTAG CAGAGAACTG ACCAGAACAT GTCTGGTCGT AAAGCTCAGG GAAAAACCCT GGGCGTCAAT 121 SGRKAQGKTLGVN ATGGTACGAC GAGGAGTTCG CTCCTTGTCA AACAAAATAA 161 MVRRGVRSLSNKI AACAAAAAC AAAACAAATT GGAAACAGAC CTGGACCTTC 201 KOKT KQI GNRP GPS 241 AAGAGGTGTT CAAGGATTTA TCTTTTTCTT TTTGTTCAAC R G V Q G F I F F F L F N ATTTTGACTG GAAAAAAGAT CACAGCCCAC CTAAAGAGGT 281 ILTG KKI TAH LKR 321 TGTGGAAAAT GCTGGACCCA AGACAAGGCT TGGCTGTTCT LWKM LDP RQGL AVL AAGGAAAGTC AAGAGAGTGG TGGCCAGTTT GATGAGAGGA R K V K R V V A S L M R G 401 TTGTCCTCAA GGAAACGCCG TTCCCATGAT GTTCTGACTG L S S R K R R S H D V L T TGCAATTCCT AATTTTGGGA ATGCTGTTGA TGACGGGTGG 441 V Q F L I L G M L L M T G G AGTTACCCTC TCTAACTTCC AAGGGAAGGT GATGATGACG 481 V T L S N F Q G K V M M T GTAAATGCTA CTGACGTCAC AGATGTCATC ACGATTCCAA 521 V N A T D V T D V I T I P CAGCTGCTGG AAAGAACCTA TGCATTGTCA GAGCAATGGA TAAG KNL CIVR AMD TGTGGGATAC ATGTGCGATG ATACTATCAC TTATGAATGC 601 V G Y M C D D T I T Y E C CCAGTGCTGT CGGCTGGTAA TGATCCAGAA GACATCGACT 641 P V L S A G N D P E D I D

681 GTTGGTGCAC AAAGTCAGCA GTCTACGTCA GGTATGGAAG CWCTKSAVYVRYGR 721 ATGCACCAAG ACACGCCACT CAAGACGCAG TCGGAGGTCA CTKTRHSRRS RRS CTGACAGTGC AGACACAGG AGAAAGCACT CTAGCGAACA 761 L T V Q. T H G E S T L A N · AGAAGGGGC TTGGATGGAC AGCACCAAGG CCACAAGGTA K K G A W M D S T K A T R Y TTTGGTAAAA ACAGAATCAT GGATCTTGAG GAACCCTGGA 841 LVKTESWILRNPG TATGCCCTGG TGGCAGCCGT CATTGGTTGG ATGCTTGGGA YALVAAVIGWMLG GCAACACCAT GCAGAGAGTT GTGTTTGTCG TGCTATTGCT 921 SNTMQRVVFVV LLL TTTGGTGGCC CCAGCTTACA GCTTCAACTG CCTTGGAATG 961 LVAPAYS FNC LG M AGCAACAGAG ACTTCTTGGA AGGAGTGTCT GGAGCAACAT 1001 SNRDFLEGVSGAT GGGTGGATTT GGTTCTCGAA GGCGACAGCT GCGTGACTAT 1041 W V D L V L E G D S C V T I CATGTCTAAG GACAAGCCTA CCATCGACGT CAAGATGATG 1081 MSK DKPT I DV K M M 1121 AATATGGAGG CGGCCAACCT GGCAGAGGTC CGCAGTTATT N M E A A N L A E V R S Y GCTATTTGGC TACCGTCAGC GATCTCTCCA CCAAAGCTGC 1161 TVSDLST CYLA ATGCCCGACC ATGGGAGAAG CTCACAATGA CAAACGTGCT 1201 C P T M G E A H N D K R A GACCCAGCTT TTGTGTGCAG ACAAGGAGTG GTGGACAGGG 1241 DPAF VCR QG V GCTGGGGCAA CGGCTGCGGA TTTTTTGGCA AAGGATCCAT 1281 G W G N G C G F F G K G S I TGACACATGC GCCAAATTTG CCTGCTCTAC CAAGGCAATA 1321 D T C A K F A C S T K A I

ChimerivaxWN02 Final Product Bottled (Run 1) L/N# 02H01; P/N# FP-0008 [Strand]

1361 GGAAGAACCA TCTTGAAAGA GAATATCAAG TACGAAGTGG GRTILKENIK YEV. · CCATTITTGT CCATGGACCA ACTACTGTGG AGTCGCACGG 1401 A I F V H G P T T V E S H G AAATTACTCC ACACAGGTTG GAGCCACTCA GGCCGGCCGA 1441 NYSTQVGATQAGR TTCAGCATCA CTCCTGCTGC GCCTTCATAC ACACTAAAGC 1481 FSIT PAA PSY TLK TTGGAGAATA TGGAGAGGTG ACAGTGGACT GTGAACCACG 1521 LGEYGEV TVDC EPR GTCAGGGATT GACACCAATG CATACTACGT GATGACTGTT 1561 SGIDTNAYYVMTV GGAACAAGA CGTTCTTGGT CCATCGTGAG TGGTTCATGG 1601 GTKTFLV HRE WFM ACCTCAACCT CCCTTGGAGC AGTGCTGGAA GTACTGTGTG 1641 DLNL PWS SAGS TVW GAGGAACAGA GAGACGTTAA TGGAGTTTGA GGAACCACAC 1681 RNRETLMEFEEPH 1721 GCCACGAAGC AGTCTGTGAT AGCATTGGGC TCACAAGAGG ATKQ SVI ALG SQE. GAGCTCTGCA TCAAGCTTTG GCTGGAGCCA TTCCTGTGGA GALH QAL AGAI PVE ATTTCAAGC AACACTGTCA AGTTGACGTC GGGTCATTTG . 1801 FSS.NTVKLTSGHL. · AAGTGTAGAG TGAAGATGGA AAAATTGCAG TTGAAGGGAA 1841 K C R V K M E K L Q L K G CAACCTATGG CGTCTGTTCA AAGGCTTTCA AGTTTCTTAG 1881 T T Y G V C S K A F K F L R GACTCCCGTG GACACCGGTC ACGGCACTGT GGTGTTGGAA 1921 TPV DTGHGTV V LE TTGCAGTACA CTGGCACGGA TGGACCTTGC AAAGTTCCTA 1961 LOYT GTD GPC KVP TCTCGTCAGT GGCTTCATTG AACGACCTAA CGCCAGTGGG 2001 ISSV ASL NDLT PVG

2041 CAGATTGGTC ACTGTCAACC CTTTTGTTTC AGTGGCCACG RLV T V N P F V S V A T GCCAACGCTA AGGTCCTGAT TGAATTGGAA CCACCCTTTG 2081 ANAK VLI ELE PPF 2121 GAGACTCATA CATAGTGGTG GGCAGAGGAG AACAACAGAT G D S Y I V V G R G E Q Q I CAATCACCAT TGGCACAAGT CTGGAAGCAG CATTGGCAAA 2161 NHH WHKS GSS I GK GCCTTTACAA CCACCCTCAA AGGAGCGCAG AGACTAGCCG 2201 AFTT TLK GAQ RLA CTCTAGGAGA CACAGCTTGG GACTTTGGAT CAGTTGGAGG ALGD TAW DFGS VGG GGTGTTCACT AGTGTTGGGC GGGCTGTCCA TCAAGTGTTC 2281 V F T S V G R A V H Q V F GGAGGAGCAT TCCGCTCACT GTTCGGAGGC ATGTCCTGGA 2321 GGAFRSLFGGMSW TAACGCAAGG ATTGCTGGGG GCTCTCCTGT TGTGGATGGG 2361 ITQG LLG ALLL W M G CATCAATGCT CGTGATAGGT CCATAGCTCT CACGTTTCTC 2401 INARDRS IAL.TFL GCAGTTGGAG GAGTTCTGCT CTTCCTCTCC GTGAACGTGG AVGGVLL FLS V N V GCGCCGATCA AGGATGCGCC ATCAACTTTG GCAAGAGAGA 2481 GADQ GCAINFG KRE GCTCAAGTGC GGAGATGGTA TCTTCATATT TAGAGACTCT 2521 LKC GDGI FIF RDS GATGACTGGC TGAACAAGTA CTCATACTAT CCAGAAGATC 2561 D D W L N K Y S Y Y P E D CTGTGAAGCT TGCATCAATA GTGAAAGCCT CTTTTGAAGA 2601 P V K L A S I V K A S F E E AGGGAAGTGT GGCCTAAATT CAGTTGACTC CCTTGAGCAT 2641 G K C G L N S V D S L E H 2681 GAGATGTGGA GAAGCAGGGC AGATGAGATC AATGCCATTT EMWR SRADEI NAI

TTGAGGAAAA CGAGGTGGAC ATTTCTGTTG TCGTGCAGGA FEENEVD ISVV V Q D TCCAAAGAAT GTTTACCAGA GAGGAACTCA TCCATTTTCC 2761 PKN VYQR GTH PFS AGAATTCGGG ATGGTCTGCA GTATGGTTGG AAGACTTGGG 2801 RIRDGLQYGWKTW GTAAGAACCT TGTGTTCTCC CCAGGGAGGA AGAATGGAAG G K N L V F S P G R K N G S 2881 CTTCATCATA GATGGAAAGT CCAGGAAAGA ATGCCCGTTT FIIDGKS RKE CPF 2921 TCAAACCGGG TCTGGAATTC TTTCCAGATA GAGGAGTTTG S N R V W N S F Q I E E F GGACGGGAGT GTTCACCACA CGCGTGTACA TGGACGCAGT 2961 G T G V F T T R V Y M D A V CTTTGAATAC ACCATAGACT GCGATGGATC TATCTTGGGT 3001 FEYTIDC DGS ILG GCAGCGGTGA ACGGAAAAAA GAGTGCCCAT GGCTCTCCAA 3041 A A V N G K K S A H G S P CATTITGGAT GGGAAGTCAT GAAGTAAATG GGACATGGAT 3081 TFWM GSH EVNG TWM GATCCACACC TTGGAGGCAT TAGATTACAA GGAGTGTGAG 3121 I H T L E A L D Y K E C E TGGCCACTGA CACATACGAT TGGAACATCA GTTGAAGAGA 3161 WPLT HTI GTS VEE. 3201 GTGAAATGTT CATGCCGAGA TCAATCGGAG GCCCAGTTAG SEMF MPR SIGG PV S CTCTCACAAT CATATCCCTG GATACAAGGT TCAGACGAAC 3241 S H N H I P G Y K V Q T N GGACCTTGGA TGCAGGTACC ACTAGAAGTG AAGAGAGAAG 3281 G P W M 'Q V P L E V K R E CTTGCCCAGG GACTAGCGTG ATCATTGATG GCAACTGTGA 3321 A C P G T S V I I D G TGGACGGGGA AAATCAACCA GATCCACCAC GGATAGCGGG 3361 GRGKSTR STT D SG

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## ChimerivaxWN02 Final Product Bottled (Run 1) L/N# 02H01; P/N# FP-0008 [Strand]

3401 AAAGTTATTC CTGAATGGTG TTGCCGCTCC TGCACAATGC K V I P E W C C R S C T M CGCCTGTGAG CTTCCATGGT AGTGATGGGT GTTGGTATCC 3441 PPVS FHG SDGC WYP CATGGAAATT AGGCCAAGGA AAACGCATGA AAGCCATCTG 3481 MEIRPRKTHE SHL 3521 GTGCGCTCCT GGGTTACAGC TGGAGAAATA CATGCTGTCC V R S W V T A G E I H A V 3561 CTTTTGGTTT GGTGAGCATG ATGATAGCAA TGGAAGTGGT PFGL VSM MIAMEVV CCTAAGGAAA AGACAGGGAC CAAAGCAAAT GTTGGTTGGA 3601 L'R'K R'QGP,KQM LVG 3641 GGAGTAGTGC TCTTGGGAGC AATGCTGGTC GGGCAAGTAA G V V L L G A M L V G Q V CTCTCCTTGA TTTGCTGAAA CTCACAGTGG CTGTGGGATT TLLD LLK LTVA VGL 3721 GCATTTCCAT GAGATGAACA ATGGAGGAGA CGCCATGTAT H F H E M N N G G D A M Y ATGGCGTTGA TTGCTGCCTT TTCAATCAGA CCAGGGCTGC 3761 MALIAAF SIR PGL TCATCGGCTT TGGGCTCAGG ACCCTATGGA GCCCTCGGGA 3801 LIGF GLR TLWS PRE ACGCCTTGTG CTGACCCTAG GAGCAGCCAT GGTGGAGATT 3841 RLV LTLG AAM VEI GCCTTGGGTG GCGTGATGGG CGGCCTGTGG AAGTATCTAA 3881 . A L G G V M G G L W K Y L ATGCAGTTTC TCTCTGCATC CTGACAATAA ATGCTGTTGC 3921 NAVS LCI LTINAVA TTCTAGGAAA GCATCAAATA CCATCTTGCC CCTCATGGCT 3961 S R K A S N T I L P L M A CTGTTGACAC CTGTCACTAT GGCTGAGGTG AGACTTGCCG 4001 LLTPVTMAEVRLA CAATGTTCTT TTGTGCCATG GTTATCATAG GGGTCCTTCA 4041 AMF.F CAM VIIG VL.H

CCAGAATTTC AAGGACACCT CCATGCAGAA GACTATACCT 4081 Q N F K D T S M Q K T I P. CTGGTGGCCC TCACACTCAC ATCTTACCTG GGCTTGACAC 4121 L V A L T L T S Y L G L T AACCTTTTT GGGCCTGTGT GCATTTCTGG CAACCCGCAT 4161 Q P F L G L C A F L A T R I ATTTGGGCGA AGGAGTATCC CAGTGAATGA GGCACTCGCA 4201 FGR RSIP VNE ALA GCAGCTGGTC TAGTGGGAGT GCTGGCAGGA CTGGCTTTTC 4241 AAGL VGV LAG LAF 4281 AGGAGATGGA GAACTTCCTT GGTCCGATTG CAGTTGGAGG QEMENFL GPIA V G G ACTCCTGATG ATGCTGGTTA GCGTGGCTGG GAGGGTGGAT 4321 ... L L M M L V S V A G R V D GGGCTAGAGC TCAAGAAGCT TGGTGAAGTT TCATGGGAAG 4361 G L E L K K L G E V S W E AGGAGGCGGA GATCAGCGGG AGTTCCGCCC GCTATGATGT 4401 EEAE ISGSSAR YDV GGCACTCAGT GAACAAGGGG AGTTCAAGCT GCTTTCTGAA 4441. ALS EQGE FKL LSE GAGAAAGTGC CATGGGACCA GGTTGTGATG ACCTCGCTGG 4481 EKVPWDQVVM TSL 4521 CCTTGGTTGG GGCTGCCCTC CATCCATTTG CTCTTCTGCT ALVG AAL HPFA LLL 4561 GGTCCTTGCT GGGTGGCTGT TTCATGTCAG GGGAGCTAGG V L A G W L F H V R G A R AGAAGTGGGG ATGTCTTGTG GGATATTCCC ACTCCTAAGA R S G D V L W D I P T P K TCATCGAGGA ATGTGAACAT CTGGAGGATG GGATTTATGG 4641 I I E E C E H L E D G I Y G CATATTCCAG TCAACCTTCT TGGGGGCCTC CCAGCGAGGA 4681 I F Q S T F L G A S Q R G GTGGGAGTGG CACAGGGAGG GGTGTTCCAC ACAATGTGGC 4721 V G V A Q G G V F H T M W

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4761 ATGTCACAAG AGGAGCTTTC CTTGTCAGGA ATGGCAAGAA HVTRGAFLVRNGKK GTTGATTCCA TCTTGGGCTT CAGTAAAGGA AGACCTTGTC 4801 LIPSWAS VKE DLV GCCTATGGTG GCTCATGGAA GTTGGAAGGC AGATGGGATG A Y G G S W K L E G R W D GAGAGGAAGA GGTCCAGTTG ATCGCGGCTG TTCCAGGAAA 4881 GEEE VQLIAAV PGK GAACGTGGTC AACGTCCAGA CAAAACCGAG CTTGTTCAAA 4921 N V V N V Q T K P S L F K GTGAGGAATG GGGGAGAAAT CGGGGCTGTC GCTCTTGACT 4961 VRNGGEIGAVALD 5001 ATCCGAGTGG CACTTCAGGA TCTCCTATTG TTAACAGGAA Y P S G T S G S P I V N R N CGGAGAGGTG ATTGGGCTGT ACGGCAATGG CATCCTTGTC 5041 GEVIGLY GNG I LV GGTGACAACT CCTTCGTGTC CGCCATATCC CAGACTGAGG 5081 G D N S F V S A I S Q T E TGAAGGAAGA AGGAAAGGAG GAGCTCCAAG AGATCCCGAC 5121 V K E E G K E E L Q E I P T AATGCTAAAG AAAGGAATGA CAACTGTCCT TGATTTTCAT 5161 MLKKGMTTVLDFH CCTGGAGCTG GGAAGACAAG ACGTTTCCTC CCACAGATCT 5201 PGAGKTR RFL PQI TGGCCGAGTG CGCACGGAGA CGCTTGCGCA CTCTTGTGTT 5241 LAECARR RLRT LVL GGCCCCCACC AGGGTTGTTC TTTCTGAAAT GAAGGAGGCT 5281 APTRVVL SEMKEA TTTCACGGCC TGGACGTGAA ATTCCACACA CAGGCTTTTT 5321 FHGL DVK FHT QAF CCGCTCACGG CAGCGGGAGA GAAGTCATTG ATGCCATGTG 5361 SAHG SGR EVID AMC CCATGCCACC CTAACTTACA GGATGTTGGA ACCAACTAGG 5401 HAT LTYR MLE PTR

5441 GTTGTTAACT GGGAAGTGAT CATTATGGAT GAAGCCCATT V V N W E V I I M D E A H 5481 TTTTGGATCC AGCCAGCATA GCCGCTAGAG GTTGGGCAGC F L D P A S I A A R G W A A GCACAGAGCT AGGGCAAATG AAAGTGCAAC AATCTTGATG 5521 HRARANE SATILM ACAGCCACAC CGCCTGGGAC TAGTGATGAA TTTCCACATT 5561 TATPPGTSDEFPH CAAATGGTGA AATAGAAGAT GTTCAAACGG ACATACCCAG 5601 SNGE I ED V Q T D I P S. TGAGCCCTGG AACACAGGGC ATGACTGGAT CCTGGCTGAC 5641 EPW NTGH DWI LAD AAAAGGCCCA CGGCATGGTT CCTTCCATCC ATCAGAGCTG 5681 KRPTAWF LPS IRA CAAATGTCAT GGCTGCCTCT TTGCGTAAGG CTGGAAAGAG 5721 . ANVM AAS LRKA GKS TGTGGTGGTC CTGAACAGGA AAACCTTTGA GAGAGAATAC 5761 V V V L N R K T F E R E Y CCCACGATAA AGCAGAAGAA ACCTGACTTT ATATTGGCCA 5801 PTIKQKK PDF I LA CTGACATAGC TGAAATGGGA GCCAACCTTT GCGTGGAGCG 5841 T D I A E M G A N L C V E R AGTGCTGGAT TGCAGGACGG.CTTTTAAGCC TGTGCTTGTG 5881 VLDCRTAFKPVLV. GATGAAGGGA GGAAGGTGGC AATAAAAGGG CCACTTCGTA 5921 DEGRKVAIKG PLR TCTCCGCATC CTCTGCTGCT CAAAGGAGGG.GGCGCATTGG 5961 ISAS SAA QRRG RIG GAGAAATCCC AACAGAGATG GAGACTCATA CTACTATTCT 6001 RNPNRDGDSYYYS GAGCCTACAA GTGAAAATAA TGCCCACCAC GTCTGCTGGT 6041 EPTSENN AHH VCW TGGAGGCCTC AATGCTCTTG GACAACATGG AGGTGAGGGG 6081 LEAS M L L D N M E V R G

ChimerivaxWN02 Final Product Bottled (Run 1) L/N# 02H01; P/N# FP-0008 [Strand]

TGGAATGGTC GCCCCACTCT ATGGCGTTGA AGGAACTAAA 6121 G M V A P L Y G V E G T K 6161 ACACCAGTTT CCCCTGGTGA AATGAGACTG AGGGATGACC TPVSPGEMRLRDD AGAGGAAAGT CTTCAGAGAA CTAGTGAGGA ATTGTGACCT Q R K V F R E L V R N C D L 6241 GCCCGTTTGG CTTTCGTGGC AAGTGGCCAA GGCTGGTTTG PVW LSWQ VAK AGL AAGACGAATG ATCGTAAGTG GTGTTTTGAA GGCCCTGAGG 6281 KTND RKW CFE GPE 6321 AACATGAGAT CTTGAATGAC AGCGGTGAAA CAGTGAAGTG EHEILND SGET VKC 6361 CAGGGCTCCT GGAGGAGCAA AGAAGCCTCT GCGCCCAAGG R A P G G A K K P L R P R 6401 TGGTGTGATG AAAGGGTGTC ATCTGACCAG AGTGCGCTGT W C D E R V S S D Q S A L 6441 CTGAATTTAT TAAGTTTGCT GAAGGTAGGA GGGGAGCTGC SEFIKFAEGRR GAA TGAAGTGCTA GTTGTGCTGA GTGAACTCCC TGATTTCCTG EVL VVLS ELP D F L GCTAAAAAG GTGGAGAGGC AATGGATACC ATCAGTGTGT 6521 AKKG GEA M D T I S V 6561 TCCTCCACTC TGAGGAAGGC TCTAGGGCTT ACCGCAATGC FLHSEEG SRAY RNA 6601 ACTATCAATG ATGCCTGAGG CAATGACAAT AGTCATGCTG LSM MPBAMTI V ML 6641 TTTATACTGG CTGGACTACT GACATCGGGA ATGGTCATCT FILAGLL TSG M V I TTTTCATGTC TCCCAAAGGC ATCAGTAGAA TGTCTATGGC 6681 FFMS PKG ISRM SMA GATGGGCACA ATGGCCGGCT GTGGATATCT CATGTTCCTT 6721 MGT MAGC GYL MFL GGAGGCGTCA-AACCCACTCA CATCTCCTAT GTCATGCTCA 6761 G G. V K P T H I S Y V M L

TATTCTTTGT CCTGATGGTG GTTGTGATCC CCGAGCCAGG 6801 I F F V L M V V V I P E P G GCAACAAAGG TCCATCCAAG ACAACCAAGT GGCATACCTC 6841 Q Q R S I Q D N Q V A Y L ATTATTGGCA TCCTGACGCT GGTTTCAGCG GTGGCAGCCA 6881 I I G I L T L V S A V A A ACGAGCTAGG CATGCTGGAG AAAACCAAAG AGGACCTCTT 6921 NELG MLE KTKE DLF TGGGAAGAAG AACTTAATTC CATCTAGTGC TTCACCCTGG 6961 G K K N L I P S S A S P W AGTTGGCCGG ATCTTGACCT GAAGCCAGGA GCTGCCTGGA 7001 SWPD LDL KPG AAW CAGTGTACGT TGGCATTGTT ACAATGCTCT CTCCAATGTT 7041 T V Y V G I V T M L S P M L GCACCACTGG ATCAAAGTCG AATATGGCAA CCTGTCTCTG 7081 H H W I K V E Y G N L S L TCTGGAATAG CCCAGTCAGC CTCAGTCCTT TCTTTCATGG 7121 S G I A Q S A S V L S F M ACAAGGGGAT ACCATTCATG AAGATGAATA TCTCGGTCAT 7161 D K G I P F M K M N I S V I AATGCTGCTG GTCAGTGGCT GGAATTCAAT AACAGTGATG 7201 M L L V S G W N S I T V M CCTCTGCTCT GTGGCATAGG GTGCGCCATG CTCCACTGGT 7241 PLLCGIGCAM LHW CTCTCATTTT ACCTGGAATC AAAGCGCAGC AGTCAAAGCT 7281 S L I L P G I K A Q Q S K L TGCACAGAGA AGGGTGTTCC ATGGCGTTGC CAAGAACCCT 7321 AQRRVFHGVAKNP GTGGTTGATG GGAATCCAAC AGTTGACATT GAGGAAGCTC 7361 V V D G N P T V D I E E A CTGAAATGCC TGCCCTTTAT GAGAAGAAAC TGGCTCTATA 7401 PEMPALYEKKL ALY TCTCCTTCTT GCTCTCAGCC TAGCTTCTGT TGCCATGTGC 7441 L L L A L S L A S V A M C

AGAACGCCCT TTTCATTGGC TGAAGGCATT GTCCTAGCAT 7481 RTPFSLAEGIVLA CAGCTGCCTT AGGGCCGCTC ATAGAGGGAA ACACCAGCCT SAAL GPL I E GN T S L TCTTTGGAAT GGACCCATGG CTGTCTCCAT GACAGGAGTC 7561 L W N G P M A V S M T G V ATGAGGGGGA ATCACTATGC TTTTGTGGGA GTCATGTACA 7601 M R G N H Y A F V G V M Y ATCTATGGAA GATGAAAACT GGACGCCGGG GGAGCGCGAA 7641 NLWKMKTGRRGSAN 7681 TGGAAAAACT TTGGGTGAAG TCTGGAAGAG GGAACTGAAT G K T L G E V W K R E L N CTGTTGGACA AGCGACAGTT TGAGTTGTAT AAAAGGACCG 7721 LLDKRQFELYKRT ACATTGTGGA GGTGGATCGT GATACGGCAC GCAGGCATTT DIVE VDR DTAR RHL GGCCGAAGGG AAGGTGGACA CCGGGGTGGC GGTCTCCAGG 7801 AEGKVDT GVA VSR GGGACCGCAA AGTTAAGGTG GTTCCATGAG CGTGGCTATG 7841 GTAKLRWFHERGY TCAAGCTGGA AGGTAGGGTG ATTGACCTGG GGTGTGGCCG 7881 V K L E G R V I D L G C G R CGGAGGCTGG TGTTACTACG CTGCTGCGCA AAAGGAAGTG . 7921 . G G W C Y Y A A A Q K E V. 7961 AGTGGGGTCA AAGGATTTAC TCTTGGAAGA GACGGCCATG S G V K G F T L G R D G H 8001 AGAAACCCAT GAATGTGCAA AGTCTGGGAT GGAACATCAT E K P M N V Q S L G W N I I CACCTTCAAG GACAAAACTG ATATCCACCG CCTAGAACCA 8041 TFK DKTD IHR LEP GTGAAATGTG ACACCCTTTT GTGTGACATT GGAGAGTCAT 8081 V K C D T L L C D I G E S CATCGTCATC GGTCACAGAG GGGGAAAGGA CCGTGAGAGT 8121 SSSS V T E G E R T V R V

TCTTGATACT GTAGAAAAAT GGCTGGCTTG TGGGGTTGAC 8161 L D T V E K W L A C G V D AACTTCTGTG TGAAGGTGTT AGCTCCATAC ATGCCAGATG N F C V K V L A P Y M P D TTCTTGAGAA ACTGGAATTG CTCCAAAGGA GGTTTGGCGG 8241 V L E K L E L L Q R R F G G AACAGTGATC AGGAACCCTC TCTCCAGGAA TTCCACTCAT 8281 TVIRNPLSRN GAAATGTACT ACGTGTCTGG AGCCCGCAGC AATGTCACAT 8321 EMYYVSGARSNVT TTACTGTGAA CCAAACATCC CGCCTCCTGA TGAGGAGAAT 8361 FTVN-QTS RLLM RRM GAGGCGTCCA ACTGGAAAAG TGACCCTGGA GGCTGACGTC 8401 RRPTGKVTLEADV ATCCTCCCAA TTGGGACACG CAGTGTTGAG ACAGACAAGG 8441 ILPIGTR SVE TDK GACCCCTGGA CAAAGAGGCC ATAGAAGAAA GGGTTGAGAG 8481 GPLD KEAIEER VER GATAAAATCT GAGTACATGA CCTCTTGGTT TTATGACAAT 8521 IKSEYMT SWF Y D N GACAACCCCT ACAGGACCTG GCACTACTGT GGCTCCTATG 8561 D N P Y R T W H Y C G S Y TCACAAAAC CTCCGGAAGT GCGGCGAGCA TGGTAAATGG 8601 V T K T S G S A A S M V, N G TGTTATTAAA ATTCTGACAT ATCCATGGGA CAGGATAGAG 8641 VIKILTY PWD RIE GAGGTCACAA GAATGGCAAT GACTGACACA ACCCCTTTTG 8681. EVTR MAM TOT TPF GACAGCAAAG AGTGTTTAAA GAAAAAGTTG ACACCAGAGC 8721 GQQRVFKEKVDTRA AAAGGATCCA CCAGCGGGAA CTAGGAAGAT CATGAAAGTT 8763 K D P P A G T R K I M K V GTCAACAGGT GGCTGTTCCG CCACCTGGCC AGAGAAAAGA 8801 VNRW LFR H L A R E K ChimerivaxWN02 Final Product Bottled (Run 1) L/N# 02H01; P/N# FP-0008 [Strand]

8841 ACCCCAGACT GTGCACAAAG GAAGAATTTA TTGCAAAAGT NPRL CTK EEFI AKV 8881 CCGAAGTCAT GCAGCCATTG GAGCTTACCT GGAAGAACAA R S H A A I G A Y L E E Q GAACAGTGGA AGACTGCCAA TGAGGCTGTC CAAGACCCAA E Q W K T A N E A V Q D P AGTTCTGGGA ACTGGTGGAT GAAGAAAGGA AGCTGCACCA 8961 K F W E L V D E E R K L H Q ACAAGGCAGG TGTCGGACTT GTGTGTACAA CATGATGGGG 9001 Q G R C R T C V Y N M M G AAAAGAGAGA AGAAGCTGTC AGAGTTTGGG AAAGCAAAGG 9041 KREKKLSEFGKAK GAAGCCGTGC CATATGGTAT ATGTGGCTGG GAGCGCGGTA 9081 G S R A I W Y M W L G A R Y TCTTGAGTTT GAGGCCCTGG GATTCCTGAA TGAGGACCAT 9121 LEFEALG FLN EDH: TGGGCTTCCA GGGAAAACTC AGGAGGAGGA GTGGAAGGCA 9161 WASRENS GGG VEG TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC 9201 IGLQYLGYVIR DLA TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT 9241 A M D G G G F Y A D D T A 9281 GGATGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG G W D T R I T E A D L D D 9321 AACAGGAGAT CTTGAACTAC ATGAGCCCAC ATCACAAAAA EQEILNY MSPH HKK 9361 ACTGGCACAA GCAGTGATGG AAATGACATA CAAGAACAAA LAQAVMEMTYKNK 9401 GTGGTGAAAG TGTTGAGACC AGCCCCAGGA GGGAAAGCCT VVKV LRP APG GKA ACATGGATGT CATAAGTCGA CGAGACCAGA GAGGATCCGG 9441 Y M D V I S R R D Q R G S G GCAGGTAGTG ACTTATGCTC TGAACACCAT CACCAACTTG 9481 Q V V T Y A L N T I T N L

ChimerivaxWN02 Final Product Bottled (Run 1) L/N# 02H01; P/N# FP-0008 [Strand]

AAAGTCCAAT TGATCAGAAT GGCAGAAGCA GAGATGGTGA K V Q L I R M A E A E M V TACATCACCA ACATGTTCAA GATTGTGATG AATCAGTTCT 9561 I H H Q H V Q D C D E S V L GACCAGGCTG GAGGCATGGC TCACTGAGCA CGGATGTGAC 9601 TRLEAWLTEH GCD AGACTGAAGA GGATGGCGGT GAGTGGAGAC GACTGTGTGG 9641 R L K R M A V S G D D C V TCCGGCCCAT CGATGACAGG TTCGGCCTGG CCCTGTCCCA 9681. V R P I D D R F G L A L S H TCTCAACGCC ATGTCCAAGG TTAGAAAGGA CATATCTGAA 9721 L N A M S K V R K D I S E TGGCAGCCAT CAAAAGGGTG GAATGATTGG GAGAATGTGC WOPSKGWNDWENV 9801 CCTTCTGTTC CCACCACTTC CATGAACTAC AGCTGAAGGA PFCS HHF HELQ L K D TGGCAGGAGG ATTGTGGTGC CTTGCCGAGA ACAGGACGAG 9841 GRRIVVPCREQDE CTCATTGGGA GAGGAAGGGT GTCTCCAGGA AACGGCTGGA 9881 L I G R G R V S P G N G W TGATCAAGGA AACAGCTTGC CTCAGCAAAG CCTATGCCAA 9921 MIKE TACLSKA YAN CATGTGGTCA CTGATGTATT TTCACAAAAG GGACATGAGG 9961 MWSLMYF.HKRDMR. 10001 CTACTGTCAT TGGCTGTTTC CTCAGCTGTT CCCACCTCAT L L S L A V S SAV PTS GGGTTCCACA AGGACGCACA ACATGGTCGA TTCATGGGAA 10041 W V P Q G R T T W S I H G K 10081 AGGGGAGTGG ATGACCACGG AAGACATGCT TGAGGTGTGG GEW M.TTE DML EVW 10121 AACAGAGTAT GGATAACCAA CAACCCACAC ATGCAGGACA NRVWITNNPHMQD 10161 AGACAATGGT GAAAAAATGG AGAGATGTCC CTTATCTAAC K T M V K K W R D V P Y L T

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10201	CAAGAGACAA GACAAGCTGT GCGGATCACT GATTGGAATG
٠.	KRQ DKLC GSLIGM
10241	ACCAATAGGG CCACCTGGGC CTCCCACATC CATTTAGTCA
	TNRATWASHIHLV
10281	TCCATCGTAT CCGAACGCTG ATTGGACAGG AGAAATACAC
	I H R I R T L I G Q E K Y T
10321	TGACTACCTA ACAGTCATGG ACAGGTATTC TGTGGATGCT
	DYL TVMD RYS VDA
10361	GACCTGCAAC TGGGTGAGCT TATCTGAAAC ACCATCTAAC
	DLQLGELI
10401	AGGAATAACC GGGATACAAA CCACGGTGG AGAACCGGAC
	THE COLD CHARLES COLD CAMPANANCO NOCCCOCCO
10441	TCCCCACAAC CTGAAACCGG GATATAAACC ACGGCTGGAG
••	
10481	AACCGGACTC CGCACTTAAA ATGAAACAGA AACCGGGATA
10521	AAAACTACGG ATGGAGAACC GGACTCCACA CATTGAGACA
`	
10561	GAAGAAGTTG TCAGCCCAGA ACCCCACACG AGTTTTGCCA
10601	CTGCTAAGCT GTGAGGCAGT GCAGGCTGGG ACAGCCGACC
10801	Clockwoor or another control of the
10641	TCCAGGTTGC GAAAAACCTG GTTTCTGGGA CCTCCCACCC
•	
10681	. CAGAGTAAAA AGAACGGAGC CTCCGCTACC ACCCTCCCAC
10721	GTGGTGGTAG AAAGACGGGG TCTAGAGGTT AGAGGAGACC
-	
10761	CTCCAGGGAA CAAATAGTGG GACCATATTG ACGCCAGGGA
10801	AAGACCGGAG TGGTTCTCTG CTTTTCCTCC AGAGGTCTGT
10043	GAGCACAGTT TGCTCAAGAA TAAGCAGACC TTTGGATGAC
10841	CAGCACAGII IGCICATORE INTOCIONCE IIIGGAIGAC

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ChimerivaxWN02 Final Product Bottled (Run 1) L/N# 02H01; P/N# FP-0008 [Strand]

10881 AAACACAAAA CCACAA

1	NGTAAATCCT GTGTGCTAAT TGAGGTGCAT TGGTCTGCAA
41	ATCGAGTTGC TAGGCAATAA ACACATTTGG ATTAATTTTA
81	ATCGTTCGTT GAGCGATTAG CAGAGAACTG ACCAGAACAT
121	GTCTGGTCGT AAAGCTCAGG GAAAAACCCT GGGCGTCAAT
	S G R K A Q G K T L G V N
161	ATGGTACGAC GAGGAGTTCG CTCCTTGTCA AACAAAATAA
201	M V R R G V R S L S N K I AACAAAAAC AAAACAAATT GGAAACAGAC CTGGACCTTC
	K Q K T K Q I G N R P G P S
241	AAGAGGTGTT CAAGGATTTA TCTTTTTCTT TTTGTTCAAC
281	R G V Q G F I F F F L F N ATTTTGACTG GAAAAAAGAT CACAGCCCAC CTAAAGAGGT
321	I L T G K K I T A H L K R TGTGGAAAAT GCTGGACCCA AGACAAGGCT TGGCTGTTCT
;	LWKM LDP RQGL AVL
361	AAGGAAAGTC AAGAGAGTGG TGGCCAGTTT GATGAGAGGA
401	R K V K R V V A S L M R G TTGTCCTCAA GGAAACGCCG TTCCCATGAT GTTCTGACTG
441	L S S R K R R S H D V L T TGCAATTCCT AATTTTGGGA ATGCTGTTGA TGACGGGTGG
	VQFLILG MLLM TGG
481	AGTTACCCTC TCTAACTTCC AAGGGAAGGT GATGATGACG
	V T L S N F Q G K V M M T
521 ·	GTAAATGCTA CTGACGTCAC AGATGTCATC ACGATTCCAA
	V N A T D V T D V I T I P
561	CAGCTGCTGG AAAGAACCTA TGCATTGTCA GAGCAATGGA
	TAAGKNL CIVR AMD
601	TGTGGGATAC ATGTGCGATG ATACTATCAC TTATGAATGC
	V G Y M C D D T I T Y E C
641	CCAGTGCTGT CGGCTGGTAA TGATCCAGAA GACATCGACT

681	GTTGGTGCAC AAAGTCAGCA GTCTACGTCA GGTATGGAAG
	C W C T K S A V Y V R Y G R
721	ATGCACCAAG ACACGCCACT CAAGACGCAG TCGGAGGTCA
	C T K T R H S R R S R R S
761	CTGACAGTGC AGACACACGG AGAAAGCACT CTAGCGAACA
	L T V Q T H G E S T L A N
801	AGAAGGGGC TTGGATGGAC AGCACCAAGG CCACAAGGTA
	K K G A W M D S T K A T R Y
841	TTTGGTAAAA ACAGAATCAT GGATCTTGAG GAACCCTGGA
	L V K T E S W I L R N P G
881	TATGCCCTGG TGGCAGCCGT CATTGGTTGG ATGCTTGGGA
• •	YALVAAVIGWMLG
921	GCAACACCAT GCAGAGAGTT GTGTTTGTCG TGCCATTGCT
	S N T M Q R V V F V V P L L
961	TTTGGTGGCC CCAGCTTACA GCTTCAACTG CCTTGGAATG
	L V A P A Y S F N C L G M
1001	AGCAACAGAG ACTTCTTGGA AGGAGTGTCT GGAGCAACAT
	SNRD FLE GVS GAT
1041	GGGTGGATTT GGTTCTCGAA GGCGACAGCT GCGTGACTAT
	WVDLVLEGDSCVTI
1081	
	M S K D K P T I D V K M M
1121 .	AATATGGAGG CGGCCAACCT GGCAGAGGTC CGCAGTTATT
	N M E A A N L A E V R S Y
1161	GCTATTTGGC TACCGTCAGC GATCTCTCCA CCAAAGCTGC
	CYLATVS DLST KAA
1201	ATGCCCGACC ATGGGAGAAG CTCACAATGA CAAACGTGCT
	C P T M G E A H N D K R A
1241	GACCCAGCTT TTGTGTGCAG ACAAGGAGTG GTGGACAGGG
	D P A F V C R Q G V V D R
1281	
	G W G N G C G F F G K G S I
1321	TGACACATGC GCCAAATTTG CCTGCTCTAC CAAGGCAATA
	D T C A K F A C S T K A I

1361	GGAAGAACCA TCTTGAAAGA GAATATCAAG TACGAAGTGG
	G R T I L K E N I K Y E V
1401	CCATTTTTGT CCATGGACCA ACTACTGTGG AGTCGCACGG
•	A I F V H G P T T V E S H G
1441	AAATTACTCC ACACAGGTTG GAGCCACTCA GGCCGGCCGA
	N Y S T Q V G A T Q A G R
1481	TTCAGCATCA CTCCTGCTGC GCCTTCATAC ACACTAAAGC
	FSIT PAA PSY TLK
1521	
•	L G E Y G E V T V D C E P R
1561	GTCAGGGATT GACACCAATG CATACTACGT GATGACTGTT
	S G I D T N A Y Y V M T V
1601	GGAACAAAGA CGTTCTTGGT CCATCGTGAG TGGTTCATGG
	G T K T F L V H R E W F M
1641 .	ACCTCAACCT CCCTTGGAGC AGTGCTGGAA GTACTGTGTG
	D L N L P W S S A G S T V W
1681	GAGGAACAGA GAGACGTTAA TGGAGTTTGA GGAACCACAC
	RNRETLMEFEEPH
1721	GCCACGAAGC AGTCTGTGAT AGCATTGGGC TCACAAGAGG
	A T K Q S V I A L G S Q E
1761	GAGCTCTGCA TCAAGCTTTG GCTGGAGCCA TTCCTGTGGA
	GALH QALAGAI PVE
1801	ATTITCAAGC AACACTGTCA AGTTGACGTC GGGTCATTTG
	FSSNTVKLTSGHL
1841	AAGTGTAGAG TGAAGATGGA AAAATTGCAG TTGAAGGGAA
	K C R V K M E K L Q L K G CAACCTATGG CGTCTGTTCA AAGGCTTTCA AGTTTCTTAG
1881	
	T T Y G V C S K A F K F L R GACTCCCGTG GACACCGGTC ACGGCACTGT GGTGTTGGAA
1921	T P V D T G H G T V V L E
2062	
1961	TTGCAGTACA CTGGCACGGA TGGACCTTGC AAAGTTCCTA
1961	

2041	CAGATTGGTC ACTGTCAACC CTTTTGTTTC AGTGGCCACG
	R L V T V N P F V S V A T
2081 -	GCCAACGCTA AGGTCCTGAT TGAATTGGAA CCACCCTTTG
	ANAKVLIELEPPF
2121	GAGACTCATA CATAGTGGTG GGCAGAGGAG AACAACAGAT
	G D S Y I V V G R G E Q Q I
2161	CAATCACCAT TGGCACAAGT CTGGAAGCAG CATTGGCAAA
	N H H W H K S G S S I G K
2201	GCCTTTACAA CCACCCTCAA AGGAGCGCAG AGACTAGCCG
	A F T T L K G A Q R L A
2241	CTCTAGGAGA CACAGCTTGG GACTTTGGAT CAGTTGGAGG
	A L G D T A W D F G S V G G
2281	GGTGTTCACT AGTGTTGGGC GGGCTGTCCA TCAAGTGTTC
	V F T S V G R A V H Q V F
2321	GGAGGAGCAT TCCGCTCACT GTTCGGAGGC ATGTCCTGGA
	GGAFRSL FGG MSW
2361	TAACGCAAGG ATTGCTGGGG GCTCTCCTGT TGTGGATGGG
•	I T Q G L L G A L L L W M G
2401	CATCAATGCT CGTGATAGGT CCATAGCTCT CACGTTTCTC
	INA RDRS IAL TFL
2441	GCAGTTGGAG GAGTTCTGCT CTTCCTCTCC GTGAACGTGG
	A V G G V L L F L S V N V
2481	GCGCCGATCA AGGATGCGCC ATCAACTTTG GCAAGAGAGA
	GADQGCAINFGKRE
2521	GCTCAAGTGC GGAGATGGTA TCTTCATATT TAGAGACTCT
	LKCGDGIFIFRDS
2561	GATGACTGGC TGAACAAGTA CTCATACTAT CCAGAAGATC
	D D W L N K Y S Y Y P E D
2601	CTGTGAAGCT TGCATCAATA GTGAAAGCCT CTTTTGAAGA
	PVKL ASIVKAS FEE
2641	AGGGAAGTGT GGCCTAAATT CAGTTGACTC CCTTGAGCAT
	G K C G L N S V D S L E H
2681	GAGATGTGGA GAAGCAGGGC AGATGAGATC AATGCCATTT
	T M W D C D A D F T' N A T

2721 .	TTGAGGAAAA CGAGGTGGAC ATTTCTGTTG TCGTGCAGGA
•	FEENEVDISVVVQD
2761	TCCAAAGAAT GTTTACCAGA GAGGAACTCA TCCATTTTCC
	PKNVYQRGTHPFS
2801	AGAATTCGGG ATGGTCTGCA GTATGGTTGG AAGACTTGGG R I R D G L Q Y G W K T W
2841	GTAAGAACCT TGTGTTCTCC CCAGGGAGGA AGAATGGAAG
	G K N L V F S P G R K N G S
2881	
	F I I D G K S R K E C P F
2921	TCAAACCGGG TCTGGAATTC TTTCCAGATA GAGGAGTTTG
	S N R V W N S F Q I E F
2961	GGACGGGAGT GTTCACCACA CGCGTGTACA TGGACGCAGT
	G T G V F T T R V Y M D A V
3001	CTTTGAATAC ACCATAGACT GCGATGGATC TATCTTGGGT
	FEYTIDC DGS ILG
	GCAGCGGTGA ACGGAAAAAA GAGTGCCCAT GGCTCTCCAA
	A A V N G K K S A H G S P CATTTTGGAT GGGAAGTCAT GAAGTAAATG GGACATGGAT
3091	T F W M G S H E V N G T W M
3121	GATCCACACC TTGGAGGCAT TAGATTACAA GGAGTGTGAG
	I H T L E A L D Y K E C E
3161	TGGCCACTGA CACATACGAT TGGAACATCA GTTGAAGAGA
	W P L T H T I G T S V E E
3201	GTGAAATGTT CATGCCGAGA TCAATCGGAG GCCCAGTTAG
	SEMF MPR SIGG P.V S
3241	CTCTCACAAT CATATCCCTG GATACAAGGT TCAGACGAAC
	CTCTCACAAT CATATCCCTG GATACAAGGT TCAGACGAAC S H N H I P G Y K V Q T N
	CTCTCACAAT CATATCCCTG GATACAAGGT TCAGACGAAC S H N H I P G Y K V Q T N GGACCTTGGA TGCAGGTACC ACTAGAAGTG AAGAGAGAAG
3281 ·	CTCTCACAAT CATATCCCTG GATACAAGGT TCAGACGAAC S H N H I P G Y K V Q T N GGACCTTGGA TGCAGGTACC ACTAGAAGTG AAGAGAGAAG G P W M Q V P L E V K R E
3281 ·	CTCTCACAAT CATATCCCTG GATACAAGGT TCAGACGAAC S H N H I P G Y K V Q T N GGACCTTGGA TGCAGGTACC ACTAGAAGTG AAGAGAGAAG G P W M Q V P L E V K R E CTTGCCCAGG GACTAGCGTG ATCATTGATG GCAACTGTGA
3281 ·	CTCTCACAAT CATATCCCTG GATACAAGGT TCAGACGAAC S H N H I P G Y K V Q T N GGACCTTGGA TGCAGGTACC ACTAGAAGTG AAGAGAGAAG G P W M Q V P L E V K R E

3401	AAAGTTATTC CTGAATGGTG TTGCCGCTCC TGCACAATGC
-	KVIPEWCCRSCTM
3441	CGCCTGTGAG CTTCCATGGT AGTGATGGGT GTTGGTATCC
	PPVSFHGSDGCWYP
3481	CATGGAAATT AGGCCAAGGA AAACGCATGA AAGCCATCTG
	MEIRPRKTHE SHL
3521	GTGCGCTCCT GGGTTACAGC TGGAGAAATA CATGCTGTCC
	V R S W V T A G E I H A V
3561	CTTTTGGTTT GGTGAGCATG ATGATAGCAA TGGAAGTGGT
	PFGL V S M M I A M E V V
3601	CCTAAGGAAA AGACAGGGAC CAAAGCAAAT GTTGGTTGGA
•	L R K R Q G P K Q M L V G
3641	GGAGTAGTGC TCTTGGGAGC AATGCTGGTC GGGCAAGTAA
	G V V L L G A M L V G Q V
3681	CTCTCCTTGA TTTGCTGAAA CTCACAGTGG CTGTGGGATT
	TLLDLLKLTVA VGL
3721	GCATTTCCAT GAGATGAACA ATGGAGGAGA CGCCATGTAT
	H F H E M N N G G D A M Y
3761	ATGGCGTTGA TTGCTGCCTT TTCAATCAGA CCAGGGCTGC
	MALIAAF SIR PGL
3801	TCATCGGCTT TGGGCTCAGG ACCCTATGGA GCCCTCGGGA
	LIGFGLRTLWSPRE
3841	ACGCCTTGTG CTGACCCTAG GAGCAGCCAT GGTGGAGATT
	R L V L T L G A A M V E I
3881	GCCTTGGGTG GCGTGATGGG CGGCCTGTGG AAGTATCTAA
	ALGG VMG GLW KYL
3921	ATGCAGTTTC TCTCTGCATC CTGACAATAA ATGCTGTTGC
	NAVS LCI LTINAVA
3961	TTCTAGGAAA GCATCAAATA CCATCTTGCC CCTCATGGCT
	S R K A S N T . I L P L M A
4001	CTGTTGACAC CTGTCACTAT GGCTGAGGTG AGACTTGCCG
	L L T P V T M A E V R L A
4041	CAATGTTCTT TTGTGCCATG GTTATCATAG GGGTCCTTCA
	AMFF CAM VIIG VLH

	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
4081	CCAGAATTTC AAGGACACCT CCATGCAGAA GACTATACCT
	Q N F K D T S M Q K T I P
4121	CTGGTGGCCC TCACACTCAC ATCTTACCTG GGCTTGACAC
•	LVAL TLT SYL GLT
4161	AACCTTTTTT GGGCCTGTGT GCATTTCTGG CAACCCGCAT
	Q P F L G L C A F L A T R I
4201.	ATTTGGGCGA AGGAGTATCC CAGTGAATGA GGCACTCGCA
	FGR RSIP V N E A L A
4241.	GCAGCTGGTC TAGTGGGAGT GCTGGCAGGA CTGGCTTTTC
	AAGL VGV LAG LAF
4281	AGGAGATGGA GAACTTCCTT GGTCCGATTG CAGTTGGAGG
•	QEME NFL GPIA V G G
4321	ACTCCTGATG ATGCTGGTTA GCGTGGCTGG GAGGGTGGAT
-	L L M M L V S V A G R V D
4361.	GGGCTAGAGC TCAAGAAGCT TGGTGAAGTT TCATGGGAAG
	G L E L K K L G E V S W E
4401	AGGAGGCGGA GATCAGCGGG AGTTCCGCCC GCTATGATGT
	EEAE ISG SSAR Y D V
4441	GGCACTCAGT GAACAAGGGG AGTTCAAGCT GCTTTCTGAA
	ALSEQGE FKL LSE
4481	GAGAAAGTGC CATGGGACCA GGTTGTGATG ACCTCGCTGG
	EKVP W D Q V V M T S L
	CCTTGGTTGG GGCTGCCCTC CATCCATTTG CTCTTCTGCT
	ALVG AAL HPFA LLL
4561	GGTCCTTGCT GGGTGGCTGT TTCATGTCAG GGGAGCTAGG
	·VLAG.WLF HVRGAR
4601	AGAAGTGGGG ATGTCTTGTG GGATATTCCC ACTCCTAAGA
	RSGD VLW DIP TPK
4641	TCATCGAGGA ATGTGAACAT CTGGAGGATG GGATTTATGG
1011	I I E E C E H L E D G I Y G
4681	CATATTCCAG TCAACCTTCT TGGGGGCCTC CCAGCGAGGA
-001	IFQSTFLGASQRG
4721	
4141.	V C V A O G G V F H T M W

	•
4761	ATGTCACAAG AGGAGCTTTC CTTGTCAGGA ATGGCAAGAA
•	H V T R G A F L V R N G K K
4801	GTTGATTCCA TCTTGGGCTT CAGTAAAGGA AGACCTTGTC
	LIPSWAS VKE DLV
4841	GCCTATGGTG GCTCATGGAA GTTGGAAGGC AGATGGGATG
	A Y G G S W K L E G R W D
4881	GAGAGGAAGA GGTCCAGTTG ATCGCGGCTG TTCCAGGAAA
	GEEE VQLIAAV PGK
4921	GAACGTGGTC AACGTCCAGA CAAAACCGAG CTTGTTCAAA
	N V V N V Q T K P S L F K
4961	GTGAGGAATG GGGGAGAAAT CGGGGCTGTC GCTCTTGACT
	V R N G G E I G A V A L D
5001	ATCCGAGTGG CACTTCAGGA TCTCCTATTG TTAACAGGAA
	Y P S G T S G S P I V N R N
5041	CGGAGAGGTG ATTGGGCTGT ACGGCAATGG CATCCTTGTC
:	GEVIGLY GNG ILV
5081	
	G D N S F V S A I S Q T E
5121	TGAAGGAAGA AGGAAAGGAG GAGCTCCAAG AGATCCCGAC
	V K E E G K E E L Q E I P T
5161	
	M L K K G M T T V L D F H
5201	CCTGGAGCTG GGAAGACAAG ACGTTTCCTC CCACAGATCT
•	PGAGKTRRFLPQI
5241	TGGCCGAGTG CGCACGGAGA CGCTTGCGCA CTCTTGTGTT
	LAECARR RLRT LVL
5281	•
	A P T R V V L S E M K E A
5321	TTTCACGGCC TGGACGTGAA ATTCCACACA CAGGCTTTTT
	F H G L D V K F H T Q A F CCGCTCACGG CAGCGGGAGA GAAGTCATTG ATGCCATGTG
5361	•
	S A H G S G R E V I D A M C CCATGCCACC CTAACTTACA GGATGTTGGA ACCAACTAGG
5401	HAT LTYR MLE PTR
	нат штік мыв Ртк

5441	
	V V N W E V I I M D E A H
5481	TTTTGGATCC AGCCAGCATA GCCGCTAGAG GTTGGGCAGC
	F L D P A S I A A R G W A A
5521	GCACAGAGCT AGGGCAAATG AAAGTGCAAC AATCTTGATG
	HRARANE SATILM
5561	ACAGCCACAC CGCCTGGGAC TAGTGATGAA TTTCCACATT
	TATPPGTSDEFPH
5601	CAAATGGTGA AATAGAAGAT GTTCAAACGG ACATACCCAG
•	S N G E I E D V Q T D I P S
5641	TGAGCCCTGG AACACAGGGC ATGACTGGAT CCTGGCTGAC
	EPWNTGHDWILAD
5681	AAAAGGCCCA CGGCATGGTT CCTTCCATCC ATCAGAGCTG
	KRPT AWF LPS IRA
5721	CAAATGTCAT GGCTGCCTCT TTGCGTAAGG CTGGAAAGAG
	ANVM AAS LRKAGKS
5761	•
	V V V L N R K T F E R E Y
5801	CCCACGATAA AGCAGAAGAA ACCTGACTTT ATATTGGCCA
	PTIKQKK PDFILA
5841	CTGACATAGC TGAAATGGGA GCCAACCTTT GCGTGGAGCG
•	T D I A E M G A N L C V E R
5881	TOTAL TOTAL CONTROL OF THE TANGE TO THE TOTAL CONTROL OF THE TOTAL CONTROL OF THE TANGE OF THE TOTAL CONTROL OF THE TANGE
•	V L D C R T A F K P V L V
5921	THE PERSON OF TH
	DEGR KVAIKG PLR
5961	
	I S A S S A A Q R R G R I G
6001	GAGAAATCCC AACAGAGATG GAGACTCATA CTACTATTCT
9001	RNPNRDG DSYYYS
6047	GAGCCTACAA GTGAAAATAA TGCCCACCAC GTCTGCTGGT
6041	EPTSENNAHH VCW
	TGGAGGCCTC AATGCTCTTG GACAACATGG AGGTGAGGGG
6081	LEAS MLL DNME VRG
	LEAS MLL DNME VRG

TGGAATGGTC GCCCCACTCT ATGGCGTTGA AGGAACTAAA 6121 G M V A P L Y G V E G T K ACACCAGTTT CCCCTGGTGA AATGAGACTG AGGGATGACC 6161 T P V S P G E M R L R D D 6201 AGAGGAAAGT CTTCAGAGAA CTAGTGAGGA ATTGTGACCT Q R K V F R E L V R N C D L GCCCGTTTGG CTTTCGTGGC AAGTGGCCAA GGCTGGTTTG 6241 . PVW LSWQ VAK AG L AAGACGAATG ATCGTAAGTG GTGTTTTGAA GGCCCTGAGG K T N D R K W C F E G P E 6321 AACATGAGAT CTTGAATGAC AGCGGTGAAA CAGTGAAGTG EHEILNDSGET VKC 6361 CAGGGCTCCT GGAGGAGCAA AGAAGCCTCT GCGCCCAAGG RAPGGAK KPL RPR 6401 TGGTGTGATG AAAGGGTGTC ATCTGACCAG AGTGCGCTGT SAL WCDERVSSDQ 6441 CTGAATTTAT TAAGTTTGCT GAAGGTAGGA GGGGAGCTGC SEFI KFA EGRR GAA TGAAGTGCTA GTTGTGCTGA GTGAACTCCC TGATTTCCTG 6481 EVLVVLS BLP D F L GCTAAAAAG GTGGAGAGGC AATGGATACC ATCAGTGTGT 6521 AKKG GEAMDT ISV 6561 TCCTCCACTC TGAGGAAGGC TCTAGGGCTT ACCGCAATGC FLHSEEGSRAY RNA ACTATCAATG ATGCCTGAGG CAATGACAAT AGTCATGCTG 6601 . L S M M P E A M T I V M L TTTATACTGG CTGGACTACT GACATCGGGA ATGGTCATCT 6641 FILA GLL TSG M V I TTTTCATGTC TCCCAAAGGC ATCAGTAGAA TGTCTATGGC 6681 F F M S P K G ISRM GATGGGCACA ATGGCCGGCT GTGGATATCT CATGTTCCTT 672İ M G T M A G C G Y L M F L GGAGGCGTCA AACCCACTCA CATCTCCTAT GTCATGCTCA 6761 G G V K P T H I S Y V M L

TATTCTTTGT CCTGATGGTG GTTGTGATCC CCGAGCCAGG 6801 I F F V L M V V V I P E P G GCAACAAAGG TCCATCCAAG ACAACCAAGT GGCATACCTC Q Q R S I Q D N Q V A Y L ATTATTGGCA TCCTGACGCT GGTTTCAGCG GTGGCAGCCA 6881 L T L V S A IIGI 6921 ACGAGCTAGG CATGCTGGAG AAAACCAAAG AGGACCTCTT NELG MLE KTKE DLF TGGGAAGAAG AACTTAATTC CATCTAGTGC TTCACCCTGG 6961 G K K N L I P S S A S P W AGTTGGCCGG ATCTTGACCT GAAGCCAGGA GCTGCCTGGA 7001 S W P D L D L K P G A A W CAGTGTACGT TGGCATTGTT ACAATGCTCT CTCCAATGTT T V Y V G I V T M L S P M L GCACCACTGG ATCAAAGTCG AATATGGCAA CCTGTCTCTG 7081 HHWIKVE YGN LSL TCTGGAATAG CCCAGTCAGC CTCAGTCCTT TCTTTCATGG 7121 SGIAQSASVL ACAAGGGGAT ACCATTCATG AAGATGAATA TCTCGGTCAT 7161 D K G I P F M K M N I S V I AATGCTGCTG GTCAGTGGCT GGAATTCAAT AACAGTGATG 7201 M L L V S G W N S I T V M CCTCTGCTCT GTGGCATAGG GTGCGCCATG CTCCACTGGT 7241 P L L C G I G CAM L H. W 7281 CTCTCATTTT ACCTGGAATC AAAGCGCAGC AGTCAAAGCT SLIL PGIKAQQ SKL TGCACAGAGA AGGGTGTTCC ATGGCGTTGC CAAGAACCCT 7321 AQR RVFH GVA KNP GTGGTTGATG GGAATCCAAC AGTTGACATT GAGGAAGCTC 7361 V V D G N P T V D I CTGAAATGCC TGCCCTTTAT GAGAAGAAAC TGGCTCTATA 7401 PEMPALY EKKLALY TCTCCTTCTT GCTCTCAGCC TAGCTTCTGT TGCCATGTGC 7441 L L L A L S L A S V A M C

7481 AGAACGCCCT TTTCATTGGC TGAAGGCATT GTCCTAGCAT RTPFSLAEGIVLA 7521 CAGCTGCCTT AGGGCCGCTC ATAGAGGGAA ACACCAGCCT SAAL GPL IEGN TSL TCTTTGGAAT GGACCCATGG CTGTCTCCAT GACAGGAGTC 7561 L W N G P M A V S M T G V ATGAGGGGGA ATCACTATGC TTTTGTGGGA GTCATGTACA 7601 MRGNHYAFVGVMY ATCTATGGAA GATGAAAACT GGACGCCGGG GGAGCGCGAA 7641 NLWKMKT GRRG SAN TGGAAAAACT TTGGGTGAAG TCTGGAAGAG GGAACTGAAT 7681 GKT LGEV WKR ELN CTGTTGGACA AGCGACAGTT TGAGTTGTAT AAAAGGACCG 7721 L.LDKRQFELYKRT ACATTGTGGA GGTGGATCGT GATACGGCAC GCAGGCATTT 7761 DIVE VDR DTARRH L 7801 GGCCGAAGGG AAGGTGGACA CCGGGGTGGC GGTCTCCAGG AEGKVDT GVA VSR 7841 GGGACCGCAA AGTTAAGGTG GTTCCATGAG CGTGGCTATG G T A K L R W F H E R G Y 7881 TCAAGCTGGA AGGTAGGGTG ATTGACCTGG GGTGTGGCCG VKLE GRV I D L G C G R 7921 CGGAGGCTGG TGTTACTACG CTGCTGCGCA AAAGGAAGTG G G W C Y Y A A A Q K E V 7961 AGTGGGGTCA AAGGATTTAC TCTTGGAAGA GACGGCCATG S G V K G F T L G R D G H AGAAACCCAT GAATGTGCAA AGTCTGGGAT GGAACATCAT 8001 EKPM NVQ SLGW NII CACCTTCAAG GACAAAACTG ATATCCACCG CCTAGAACCA 8041 TFK DKTD IHR LEP 8081 GTGAAATGTG ACACCCTTTT GTGTGACATT GGAGAGTCAT V K C D T L L C D I G E S CATCGTCATC GGTCACAGAG GGGGAAAGGA CCGTGAGAGT 8121 SSSS V T E G E R T V R V

8161	TCTTGATACT GTAGAAAAAT GGCTGGCTTG TGGGGTTGAC
•	LDTVEKWLACGVD
8201	AACTTCTGTG TGAAGGTGTT AGCTCCATAC ATGCCAGATG
	N F C V K V L A P Y M P D
8241	TTCTTGAGAA ACTGGAATTG CTCCAAAGGA GGTTTGGCGG
	V L E K L E L L Q R R F G G
6281	AACAGTGATC AGGAACCCTC TCTCCAGGAA TTCCACTCAT
	T V I R N P L S R N S T H
8321	GAAATGTACT ACGTGTCTGG AGCCCGCAGC AATGTCACAT
	EMYYVSGARSNVT
8361	TTACTGTGAA CCAAACATCC CGCCTCCTGA TGAGGAGAAT
•	FTVNQTSRLLMRRM
8401	GAGGCGTCCA ACTGGAAAAG TGACCCTGGA GGCTGACGTC
•	RRPTGKVTLEADV
8441	ATCCTCCCAA TTGGGACACG CAGTGTTGAG ACAGACAAGG
	ILPIGTR SVE TDK
8481	GACCCCTGGA CAAAGAGGCC ATAGAAGAAA GGGTTGAGAG
	GPLDKEAIEERVER
8521	GATAAAATCT GAGTACATGA CCTCTTGGTT TTATGACAAT
	I K S E Y M T S W F Y D N
	GACAACCCCT ACAGGACCTG GCACTACTGT GGCTCCTATG
	DNPYRTWHYCGSY
8601	TCACAAAAAC CTCCGGAAGT GCGGCGAGCA TGGTAAATGG
	V T K T S G S A A S M V N G
8641	TGTTATTAAA ATTCTGACAT ATCCATGGGA CAGGATAGAG
	VIKILTYPWDRIE
8681	GAGGTCACAA GAATGGCAAT GACTGACACA ACCCCTTTTG
	EVTR M A M T D T T P F
8721	GACAGCAAAG AGTGTTTAAA GAAAAAGTTG ACACCAGAGC
	GQQRVFKEKVDTRA
8761	AAAGGATCCA CCAGCGGGAA CTAGGAAGAT CATGAAAGTT
	K D P P A G T R K I M K V
8801	GTCAACAGGT GGCTGTTCCG CCACCTGGCC AGAGAAAAGA
	TATEM LER HILAREK

	ACCCCAGACT GTGCACAAAG GAAGAATTTA TTGCAAAAGT
8841	NPRLCTKEEFIAKV
8881	
	R S H A A I G A Y L E E Q
8921	•
	EQWKTANEAVQDP
8961	
· ·	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
9001	ACAAGGCAGG TGTCGGACTT GTGTGTACAA CATGATGGGG
•	Q G R C R T C V Y N M M G
9041	AAAAGAGAGA AGAAGCTGTC AGAGTTTGGG AAAGCAAAGG
	KREK KLS EFG KAK
9081	GAAGCCGTGC CATATGGTAT ATGTGGCTGG GAGCGCGGTA
•	G S R A I W Y M W L G A R Y
9121	TCTTGAGTTT GAGGCCCTGG GATTCCTGAA TGAGGACCAT
	LEFEALG FLN EDH
9161	TGGGCTTCCA GGGAAAACTC AGGAGGAGGA GTGGAAGGCA
7202	
3101	
	WASRENS GGG VEG
	W A S R E N S G G G V E G TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC
9201	W A S R E N S G G G V E G TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC I G L Q Y L G Y V I R D L A
	W A S R E N S G G G V E G TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC I G L Q Y L G Y V I R D L A TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT
9201	WASRENS GGGVEG TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC IGLQYLGYVIR DLA TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT AMDGGGGGFYADDTA
9201	WASRENS GGGVEG TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC IGLQYLGYVIRDLA TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT AMDGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG
9201 9241 9281	WASRENS GGGVEGGC TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC IGLQYLGYVIR DLA TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT AMDGGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG GWDTRITEA
9201 9241 9281	WASRENS GGGVEG  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIR DLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITEACTAC ATGAGCCCAC ATCACAAAAA
9201 9241 9281 9321	WASRENS GGGVEGGC  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIR DLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITEACTAC ATGAGCCCAC ATCACAAAAA  EQEILNYMSPHHKKK
9201 9241 9281	WASRENS GGGVEG  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIR DLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITE ADLDD  AACAGGAGAT CTTGAACTAC ATGAGCCCAC ATCACAAAAA  EQEILNYMSPHHKKK  ACTGGCACAA GCAGTGATGG AAATGACATA CAAGAACAAA
9201 9241 9281 9321 9361	WASRENS GGGVEGGC  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIR DLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITEACTAC AGAGGCAGAC CTTGATGATG  AACAGGAGAT CTTGAACTAC ATGAGCCCAC ATCACAAAAA  EQEILNYMSPHHKKK  ACTGGCACAA GCAGTGATGG AAATGACATA CAAGAACAAA  LAQAVMEMTYKNKK
9201 9241 9281 9321	WASRENS GGGVEG  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIRDLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITEACTAC ATGAGCCCAC ATCACAAAAA  EQEILNYMSPHHKKK  ACTGGCACAA GCAGTGATGG AAATGACATA CAAGAACAAA  LAQAVMEMTYKKI
9201 9241 9281 9321 9361	WASRENS GGGVEG  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIR DLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITEACTAC ATGAGCCCAC ATCACAAAAA  EQEILNYMSPHHKKK  ACTGGCACAA GCAGTGATGG AAATGACATA CAAGAACAAA  LAQAVMEMTYKKNK  GTGGTGAAAG TGTTGAGACC AGCCCCAGGA GGGAAAGCCT  VVKVLRPAPGGKA
9201 9241 9281 9321 9361	WASRENS GGGVEG  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIRDLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGFYADDDTA  GGATGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITEAD LDD  AACAGGAGAT CTTGAACTAC ATGAGCCCAC ATCACAAAAA  EQEILNYMSPHHKKK  ACTGGCACAA GCAGTGATGG AAATGACATA CAAGAACAAA  LAQAVMEMTYKKKKKKKKKKKKKKKKKKKKKKKKKKKKKKKKKKK
9201 9241 9281 9321 9361 9401	WASRENS GGGVEG  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIRDLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITEACTAC ATGAGCCCAC ATCACAAAAA  EQEILNYMSPHHKKK  ACTGGCACAA GCAGTGATGG AAATGACATA CAAGAACAAA  LAQAVMEMTYKKNK  GTGGTGAAAG TGTTGAGACC AGCCCCAGGA GGGAAAGCCT  VVKVLRPA PGGKA  ACATGGATGT CATAAGTCGA CGAGACCAGA GAGGATCCGG
9201 9241 9281 9321 9361	WASRENS GGGVEG  TTGGCTTACA ATACCTAGGA TATGTGATCA GAGACCTGGC  IGLQYLGYVIRDLA  TGCAATGGAT GGTGGTGGAT TCTACGCGGA TGACACCGCT  AMDGGGGACA CGCGCATCAC AGAGGCAGAC CTTGATGATG  GWDTRITEACTAC ATGAGCCCAC ATCACAAAAA  EQEILNYMSPHHKKK  ACTGGCACAA GCAGTGATGG AAATGACATA CAAGAACAAA  LAQAVMEMTYKKNK  GTGGTGAAAG TGTTGAGACC AGCCCCAGGA GGGAAAGCCT  VVKVLRPA PGGKA  ACATGGATGT CATAAGTCGA CGAGACCAGA GAGGATCCGG

9521	AAAGTCCAAT TGATCAGAAT GGCAGAAGCA GAGATGGTGA
	K V Q L I R M A E A E M V
9561	TACATCACCA ACATGTTCAA GATTGTGATG AATCAGTTCT
	I H H Q H V Q D C D E S V L
9601	GACCAGGCTG GAGGCATGGC TCACTGAGCA CGGATGTGAC
	TRLEAWLTEH GCD
9641	AGACTGAAGA GGATGGCGGT GAGTGGAGAC GACTGTGTGG
	R L K R M A V S G D D C V
9681	TCCGGCCCAT CGATGACAGG TTCGGCCTGG CCCTGTCCCA
	$ \begin{array}{cccccccccccccccccccccccccccccccccccc$
9721	TCTCAACGCC ATGTCCAAGG TTAGAAAGGA CATATCTGAA
	L N A M S K V R K D I S E
9761	TGGCAGCCAT CAAAAGGGTG GAATGATTGG GAGAATGTGC
	W Q P S K G W N D W E N V
9801	CCTTCTGTTC CCACCACTTC CATGAACTAC AGCTGAAGGA
	PFCSHHFHELQLKD
9841	TGGCAGGAGG ATTGTGGTGC CTTGCCGAGA ACAGGACGAG
	GRRIVVPCREQDE
9881	CTCATTGGGA GAGGAAGGGT GTCTCCAGGA AACGGCTGGA
	LIGR GRV SPG NGW
9921	TGATCAAGGA AACAGCTTGC CTCAGCAAAG CCTATGCCAA
	MIKE TACLSKA YAN
9961	CATGTGGTCA CTGATGTATT TTCACAAAAG GGACATGAGG
	M W S L M Y F H K R D M R
10001	
	L L S L A V S S A V P T S
10041	GGGTTCCACA AGGACGCACA ACATGGTCGA TTCATGGGAA
	W V P Q G R T T W S I H G K
10081	AGGGGAGTGG ATGACCACGG AAGACATGCT TGAGGTGTGG
	GEW MTTE DML EVW
10121	AACAGAGTAT GGATAACCAA CAACCCACAC ATGCAGGACA
	NRVWITN NPH MQD
10161	AGACAATGGT GAAAAAATGG AGAGATGTCC CTTATCTAAC
•	KTMV KKW RDVP YLT

10201	CAAGAGACAA GACAAGCTGT GCGGATCACT GATTGGAATG
	K R Q D K L C G S L I G M
10241 -	ACCAATAGGG CCACCTGGGC CTCCCACATC CATTTAGTCA
	TNRATWASHI HLV
•	
10281	TCCATCGTAT CCGAACGCTG ATTGGACAGG AGAAATACAC
	I H R I R T L I G Q E K Y T
10321	TGACTACCTA ACAGTCATGG ACAGGTATTC TGTGGATGCT
	DYL TVMD RYS V D A
20367	GACCTGCAAC TGGGTGAGCT TATCTGAAAC ACCATCTAAC
10007	D L Q L G E L I
10401	AGGAATAACC GGGATACAAA CCACGGTGG AGAACCGGAC
10441	TCCCCACAAC CTGAAACCGG GATATAAACC ACGGCTGGAG
•	
10481	AACCGGACTC CGCACTTAAA ATGAAACAGA AACCGGGATA
	AAAACTACGG ATGGAGAACC GGACTCCACA CATTGAGACA
10521	WWWCIWCO WIGOWOWCC GONCIOCHEN CITTORING
•	
10561	GAAGAAGTTG TCAGCCCAGA ACCCCACACG AGTTTTGCCA
10601	CTGCTAAGCT GTGAGGCAGT GCAGGCTGGG ACAGCCGACC
	c ·
10641	TCCAGGTTGC GAAAAACCTG GTTTCTGGGA CCTCCCACCC
20012	
	CAGAGTAAAA AGAACGGAGC CTCCGCTACC ACCCTCCCAC
10681 .	CAGAGIAAAA AGAACGGAGC CICCGCIACC ACCCICCAC
10721	GTGGTGGTAG AAAGACGGGG TCTAGAGGTT AGAGGAGACC
:	
10761	CTCCAGGGAA CAAATAGTGG GACCATATTG ACGCCAGGGA
30003	AAGACCGGAG TGGTTCTCTG CTTTTCCTCC AGAGGTCTGT
TOPOT	Monocodno 1001101010 C111100100 Monodicidi
10841	GAGCACAGTT TGCTCAAGAA TAAGCAGACC TTTGGATGAC

## Chimerivax WN02 M66 variant [Strand]

10881 AAACACAAAA CCACAA

#### ### DNA Strider\*\* 1.317 ###

#### WN 02 x M66 Variant ⇒ DNA Alignment

DNA sequence 10896 bp \*GTAAATCCTGT ... ACAAAACCACAA linear
DNA sequence 10896 bp \*GTAAATCCTGT ... ACAAAACCACAA linear

Layout: Compacted
Hethod: Blocks (Martinez)
Mismatch penalty: Smaller (1)
Gap penalty: Medium (2)
Translation: Off

1	*GTAAATCLTGTGTGCTAATTGAGGTGCATTAATTTTA	
	ATCGTTCGTTCAGCGATTAGCAGAGAACTGACCAGAACATGTCTCGGTCGTAAAGCTCAGGGAAAAACCCTGGGCGTCAAT	
161	ATGGTACGACGAGGTTCGCTCCTTGTCAAACAAAATAAAACAAAAACAAAACAAATTGGAAACAGACCTGGACCTTC	240
241 241	AAGAGGTGTTCAACGATTTATCTTTTTTTTTTTTTTTTT	320 320
321	TGTGGAAAATGCTGGACCCAAGACAAGGCTTGGCTGTTCTAAGGAAAGTCAAGAGAGTGGTGGCCCAGTTTGATGAGAGGA	400
401	TTGTCCTCAAGGAAACGCCGTTCCCATGATGTTCTGACTGTGCAATTCCTAATTTTGGGAATGCTGTTGATGACGGTTGG	480
481	AGTTACCCTCTCTAACTTCCAAGGGAAGGTGATGACGGTAAATGCTACTGACGTCACAGATGTCATCACGATTCCAA	560
561	CAGCTGCTGGAAAGAACCTATGCATTGTCAGAGCAATGGATGTGGGATACATGTGCGATGATACTATCACTTATGAATGC	640
641	CCAGTGCTGTCGCCTGGTAATGATCCAGAAGACATCGACTGTTGGTGCACAAAGTCAGCAGTCTACGTCAGGTATGGAAG	720
721 721	ATGCACCAAGACACGCCACTCAAGACGCACTCGGAGGTCACTGACAGTGCAGACACACGGAGAAAGCACTCTAGCGAACA	800 800
801 801	AGAAGGGGGCTTGGATGGACAGCACCAAGGCTATTTGGTAAAAACAGAATCATGGATCTTGAGGAACCCTGGA	000
881	TATGCCCTGGTGGCAGCCGTCATTGGTTGGATGCTTGGGACCAACACCATGCAGAGAGTTGTGTTTGTCGTGCTATTGCT	960
961 961	TTTGGTGGCCCCAGCTTACAGCTTCAACTGCCTTGGAATGAGCCAACAGAGACTTCTTGGAAGGAGTGTCTGGAGCAACAT GGGTGGATTTGGTTCTCGAAGGCGACAGCTGCGTGACTATCATGTCTAAGGACAAGCCTACCATCGACGTCAAGATGATG	1040 1040
1121	AATATGGAGGGGCCAACCTGGCAGAGGTCCGCAGTTATTGCTATTTGGCTACCGTCAGCGATCTCTCCACCAAAGCTGC	1200
1201 1201	ATGCCCGACCATGGGACAAGCTCACAATGACAAACGTGCTGACCCAGCTTTTGTGTGCAGACAAGGACTGGTGGACAAGGA	1280 1280
1281	GCTGGGGCAACGCTGCCGCTTTTTTGGCAAAGGATCCATTGACACACGCCAAATTTGCCTGCTCTACCAAGGCAATA	1360 1360
1361 1361	${\tt GGAAGAACCATCTTGAAAGAGAATATCAAGTACGAAGTGGCCATTTTTGTCCATGGACCAACTACTGTGGAGTCGCACGG}$	1440

#### WN 02 x M66 Variant ⇒ DNA Alignment

1441 1441	AAATTACTCCACACGGTTGGAGCCACTCAGGCCGGCCGATTCAGCATCACTCCTGCTGCGCCTTCATACACACTAAAGC	1520 1520
1521 1521	TTGGAGAATATGGAGAGGTGACAGTGGACTGTGAACCACGGTCAGGGATTGACACCAATGCATACTACGTGATGACTGTT	1600 1600
1601 1601	GGAACAAAGACGTTCTTGGTCCATCGTGAGTGGTTCATGGACCTCAACCTCCCTTGGAGCAGTGCTGGAAGTACTGTGTG	1680 1680
1681 1681	GAGGAACAGAGAGACGTTAATGGAGTTTGAGGAACCACACGCCACGAAGCAGTCTGTGATAGCATTGGGCTCACAAGAGG	1760 1760
	GAGCTCTGCATCAAGCTTTGGCTGGAGCCATTCCTGTGGAATTTTCAAGCAACACTGTCAAGTTGACGTCGGGTCATTTG	1840 1840
	AAGTGTAGAGTGAAGATGGAAAAATTGCAGTTGAAGGGAACAACCTATGGCGTCTGTTCAAAGGCTTTCAAGTTTCTTAG	1920 1920
	GACTCCCGTGGACACCGGTCACGGCACTGTGGTGTTGGAATTGCAGTACACTGGCACGGATGGACCTTGCAAAGTTCCTA	2000 2000
2001	TCTCGTCAGTGGCTTCATTGAACGACCTAACGCCAGTGGGCAGATTGGTCACTGTCAACCCTTTTGTTTCAGTGGCCACG	2080 2080
2081 2081	GCCAACGCTAAGGTCCTGATTGAATTGGAACCACCCTTTGGAGACTCATACATA	2160 2160
	CAATCACCATTGGCACAAGTCTGGAAGCAGCATTGGCAAAGCCTTTACAACCACCCTCAAAGGAGCGCAGAGACTAGCCG	2240 2240
2241 2241	CTCTAGGAGACACAGCTTGGGACTTTGGATCAGTTGGAGGGGTGTTCACTAGTGTTGGGGGGGCTGTCCATCAAGTGTTC	2320 2320
2321 2321	GGAGGACCATTCCCCTCACTGTTCCGAGGCATGTCCTCGATAACGCAAGGATTGCTCGGGGCTCTCCTGTTGTGGATGGG	2400 2400
2401 2401	CATCAATCCTCGTGATAGGTCCATAGCTCTCACGTTTCTCGCAGTTGGAGGAGTTCTGCTCTCCTCTCCGTGAACGTGG	2480 2480
2481 2481	GCCCCGATCAAGGATGCCCCATCAACTTTGGCAAGAGAGAG	2560 2560
2561 2561	GATGACTGGCTGAACAAGTACTCATACTATCCAGAAGATCCTGTGAAGCTTGCATCAATAGTGAAAGCCTCTTTTGAAGA	2640 2640
2641 2641	AGGGAAGTGTGGCCTAAATTCAGTTGACTCCCTTGAGCATGAGATGTGGAGAAGCAGGGCAGATGAGATCAATGCCATTT	2720 2720
	TTGAGGAAAACGAGGTGGACATTTCTGTTGTCGTGCAGGATCCAAAGAATGTTTACCAGAGGAACTCATCCATTTTCC	2800 2800
2801 2801	AGAATTCGCGATGCTCTGCAGTATGGTTGGAAGACTTGGGGTAAGAACCTTGTGTTCTCCCCAGGGAGGAAGAATGGAAG	2880 2880
2881 2881	CTTCATCATAGATAGAAAGTCCAGGAAAGAATGCCCCTTTTCAAACCGGGTCTGGAATTCTTTCCAGATAGAGGAGTTTG	2960 2960
2961 2961	GGACGGGACTGTTCACCACACGCCTGTACATGGACGCAGTCTTTGAATACACCATAGACTGCGATGGATCTATCT	3040 3040
3041 3041	GCACCGCTGAACCGAAAAAAGAGTGCCCCATGGCTCTCCAACATTTTGGATGGGAAGTCATGAAGTAAATGGGACATGGAT	3120 3120
3121	GATCCACACCTTGGAGGCATTAGATTACAAGGAGTGTGAGTGGCCACTGACACATACGATTGGAACATCAGTTGAAGAGA	3200

7	102 x	M66 Variant => DNA Alignment	
	3201 3201	GTGAAATGTTCATGCCGAGATCAATCGGAGGCCCAGTTAGCTCTCACAATCATATCCCTGGATACAAGGTTCAGACGAAC	3280 3280
	3281 3281	GGACCTTGGATGCAGGTACCACTAGAAGTGAAGAGAGAGA	3360 3360
		TGGACGGGGAAAATCAACCAGATCCACCACGGATAGCGGGAAAGTTATTCCTGAATGGTGTTGCCGCTCCTGCACAATGC	3440 3440
	3441 3441	CCCCTGTGAGCTTCCATGGTAGTGATCGGTGTTGGTATCCCATGGAAATTAGGCCAAGGAAAACGCATGAAAGCCATCTG	3520 3520
	3521	GTGCGCTCCTGGGTTACAGCTGGAGAAATACATGCTGTCCCTTTTGGTTTGGTGAGCATGATGATAGCAATGGAAGTGGT	3600
	3601	CCTAAGGAAAAGACAGGGACCAAATGTTGGTTGGAGGAGTAGTGCTCTTGGGAGCAATGCTGGTCGGGCAAGTAA	3680
	3681		3760
	3761	ATGGCGTTGATTGCTGCCTTTTCAATCAGACCAGGGCTGCTCATCGGCTTTGGGCTCAGGACCCTATGGAGCCCTCGGGA	. 3840
	3841	ACCCCTTCTCCTGACCCTAGGAGCACCCATGGTGGAGATTGCCTTGGGTGGCGTGATGGGCGCCTGTGGAAGTATCTAA	3920
	3921	ATGCAGTTTCTCTGCATCCTGACAATAAATGCTGTTGCTTCTAGGAAAGCATCAAATACCATCTTGCCCCTCATGGCT	4000
•	4001	CTGTTGACACCTGTCACTATGGCTGAGGTGAGACTTGCCGCAATGTTCTTTTGTGCCATGGTTATCATAGGGGTCCTTCA	4080
	4081	CCAGAATTTCAAGGACACCTCCATGCAGAAGACTATACCTCTGGTGGCCCTCACACTCACACTCTTACCTGGGCTTGACAC	4160
	4161		. 4240
	4241	CCAGCTGGTCTAGTGGGAGTGCTGGCAGGACTGGCTTTTCAGGAGAACTTCCTTGGTCCGATTGCAGTTGGAGA	. 4320
	4321	ACTOCTGATGATGCTGGTTAGCGTGGCTGGGAGGTGGATGGGCTGAAGAAGCTTGGTGAAGTTTCATGGGAA	. 4400
	4401	AGGAGGCGGAGATCAGCGGGAGTTCCGCCCGCTATGATGTGGCACTCAGTGAACAAGGGGGAGTTCAAGCTGCTTTCTGA	. 4480
	4481	GAGAAAGTGCCATGGGACCAGGTTGTGATGACCTCGCTGGCCTTGGTTGG	. 4560
	4561	GGTCCTTCCTGGGTGGTGTTTCATGTCAGGGGAGCTAGGAGAAGTGGGGGATGTCTTGTGGGATATTCCCACTCCTAAG	. 4640
	4641	TCATCGAGGAATGTGAACATCTGGAGGATGGGATTTATGGCATATTCCAGTCAACCTTCTTGGGGGCCCTCCCAGCGAGG	. 4720
	4721	CTGGGAGTGGCACAGGGAGGGTGTTCCACACATGTGGCATGTCACAAGAGGAGCTTTCCTTGTCAGGAATGGCAAGA GTTGATTCCATCTTGGGCTTCAGTAAAGGAAGACCTTGTCGCCTATGGTGGCTCATGGAAGTTGGAAGGCAGATGGGAT	. 4800
	4801		. 4880
	4881	GALAGGAALADGICABIIGATOGGGCIGIICCAGGAAAAAACGIGGICAACGACCAGAAAAAAACGIGGICAA	. 4960

N 02 x	M66 Variant ⇒ DNA Alignment		
	GTGAGGAATGGGGGAGAAATGGGGGCTGTCGCTCTTGACTATCGGAGTGGCACTTCAGGATCTCCTATTGTTAACAGGAA	5040 5040	÷
	CGGAGAGGTGATTGGGCTGTACGGCAATGGCATCCTTGTCGGTGACAACTCCTTCGTGTCCGCCATATCCCAGACTGAGG	5120 5120	
	TGAAGGAAGGAAAGGAGGAGCTOCAAGAGATOCCGACAATGCTAAAGAAAGGAATGACAACTGTCCTTGATTTTCAT	5200 5200	١
5201	CCTGGAGCTGGGAAGACAAGACGTTTCCTCCCACAGATCTTGGCCGAGTGCGCACGGAGACGCTTGCGCACTCTTGTGTT	5280	)
5281	GGCCCCACCAGGGTTGTTCTTTCTGAAATGAAGGAGGCTTTTCACGGCCTGGACGTGAAATTCCACACACA	5360	)
5361	CCGCTCACGGCAGGAGAAGTCATTGATGCCATGTCCCATGCCACCTAACTTACAGGATGTTGGAACCAACTAGG	5440	)
5441		5520	)
5521	GCACAGAGCTAGGGCAAATGAAAGTGCAACAATCTTGATGACAGCCACACCGCCTGGGACTAGTGATGAATTTCCACATT	5600	)
5601	CAAATGCTGAAATAGAAGATGTTCAAACGGACATACCCAGTGACCCCTGGAACACAGGGCATGACTGGATCCTGGCTGAC	568	0
5681	AAAAGGCCCACGGCATGGTTCCTTCCATCAGAGCTGCAAATGTCATGGCTGCCTCTTTGCGTAAGGCTGGAAAGAG	5760	0
5761	TGTGGTGGTCCTGAACAGGAAAACCTTTGAGAGAGAATACCCCACGATAAAGCAGAAGAAACCTGACTTTATATTGGCCA	584	0.
5841	CTGACATAGCTGAAATGGGAGCCAACCTTTGCGTGGAGCGAGTGCTGGATTGCAGGACGGCTTTTAAGCCTGTGCTTGTC	592	0
5921	GATGAAGGGAGGAGGTGGCAATAAAAGGGCCACTTCGTATCTCCCCATCCTCTCCTCCTCAAAGGAGGGGGCCCATTGC	600	0
6001	GAGAAATCOCAACAGAGATGGAGACTCATACTATTCTGAGOCTACAAGTGAAAATAATGCOCACCACGACTCTGCTGGT	608	0
6081	TGCAGGCCTCAATGCTCTTGGACAACATGGAGGTGAGGGGGGGG	616	0
6161	ACACCAGTTTCCCCTGGTGAAATGAGACTGAGGGATGACCAGAGGAAAGTCTTCAGAGAACTAGTGAGGAATTGTGACC	. 624	0
6241	GCCCGTTTGGCTTTCGTGGCAAGTGGCCAAGGCTGGTTTGAAGACGAATGATCGTAAGTGGTGTTTTGAAGGCCCTGAG	. 632	0
6321	AACATGAGATCTTGAATGACAGGGGTGAAACAGTGAAGTGCAGGGCTCCTGGAGGAGGAGGAAGAAGCCTCTGGGGCCCAAG	. 640	0
6401	TGGTGTGATGAAAGGGTGTCATCTGACCAGAGTGCGCTGTCTGAATTTATTAAGTTTGCTGAAGGTAGGAGGGGAGCTG	. 648	30
6481	TGAAGTGCTAGTTGTGCTGAGTGAACTCCCTGATTTCCTGGCTAAAAAAGGTGGAGAGGCAATGGATACCATCAGTGTG	. 656	0
6561	TCCTCCACTCTGAGGAAGGCTCTAGGGCTTACCGCAATGCACTATCAATGATGCCTGAGGCAATGACAATAGTCATGCT	. 664	10
	TTTATACTGGCTGGACTACTGACATCGGGAATGGTCATCTTTTTCATGTCTCCCAAAGGCATCAGTAGAATGTCTATGG		

N	02 x	M66 Variant ⇒ DNA Alignment	
_	721 721	CATGGGCACAATGGCCGGCTGTGGATATCTCATGTTCCTTGGAGGCGTCAAACCCACTCACATCTCCTATGTCATGCTCA	6800
	5801 5801	TATTCTTTGTCCTGATGGTGGTGATCCCCGAGCCAGGCCAACAAAGGTCCATCCA	6880 6880
	5881 5881	ATTATTGGCATCCTGACGCTGGTTTCAGCGGTGGCAGCCAACGAGCTAGGCATGCTGGAGAAAACCAAAGAGGACCTCTT	6960 6960
	6961 5 <b>961</b>	${\tt TGGGAAGAAGAACTTAATTGCATCTAGTGCTTCACCCTGGAGTTGGGCGGATCTTGACCTGAAGCCAGGAGCTGGCTG$	7040 7040
	7041 7041	CAGTGTACGTTGCCATTGTTACAATGCTCTCCCAATGTTGCACCACTGGATCAAAGTCGAATATGGCAACCTGTCTCTG	7120 7120
		TCTGGAATAGCCCAGTCAGCCTCAGTCCTTTCTTCATGGACAAGGGGGATACCATTCATGAAGATGAATATCTCGGTCAT	7200 7200
	, 7201 7201	NATICETCETCACTGGCAATTCAATAACAGTGATGCCTCTGCTCTG	7280 7280
	7281 7281	CTCTCATTTTACCTGGAATCAAAGCGCAGCAGTCAAAGCTTGCACAGAGAAGGGTGTTCCATGGCGTTGCCAAGAACCCT	7360 7360
	7361 7361	CTCGTTGATCGGAATCCAACAGTTGACATTGACGAAGCTCCTGAAATGCCTGCC	7440 7440
	7441 7441	TCTCCTTCTTGCTCTCAGCCTAGCTTCTGTTGCCATGTGCAGAACGCCCTTTTCATTGGCTGAAGGCATTGTCCTAGCAT	7520 7520
	7521 7521	CASCTGCCTTAGGGCCGCTCATAGAGGGAAACACCAGCCTTCTTTGGAATGGACCCATGGCTGTCTCCATGACAGGAGTC	7600 7600
		ATCAGGGGGAATCACTATGCTTTTGTGGGAGTCATGTACAATCTATGGAAGATGAAAACTGGACGCCGGGGGGGG	7680 7680
	7681 7681	TGGAAAAACTTTGGGTGAAGTCTGGAAGAGGGAACTGAATCTGTTGGACAAGCGACAGTTTGAGTTGTATAAAAGGACCG	7760 7760
	7761 7761	ACATTGTGGAGGTGGATCGTGATACGGCACGCAGGCATTTGGCCGAAGGGGAAGGTGGACACCGGGGTGGCGGTCTCCAGG	7840 7840
	7841 7841	CGCACCCCAAACTTAAGGTGGTTCCATGAGCGTGGCTATGTCAAGCTGGAAGGTAGGGTGATTGACCTGGGGTGTGGCCC	7920 7920
	7921 7921	CCCAGGCTCGTGTTACTACCCTCCTCCCCAAAAGGAAGTGAGTG	8000 8000
		AGAAACCCATGAATGTGCAAAGTCTGGGATGGAACATCATCACCTTCAAGGACAAAACTGATATCCACCGCCTAGAACCI	. 8080 8080
	8081 8081	GTGAAATGTGACACCCTTTTGTGTGACATTGGAGAGTCATCATCGTCATCGGTCACAGAGGGGGAAAGGACCGTGAGAG	r 8160 8160
	8161 8161	TCTTGATACTGTAGAAAAATGGCTGGCTTGTGGGGTTGACAACTTCTGTGTGAAGGTGTTAGCTCCATACATGCCAGAT	8240 8240
	8241 8241	TTCTTGAGAAACTGGAATTGCTCCAAAGGAGGTTTGGCGGAACAGTGATCAGGAACCCTCTCTCCAGGAATTCCACTCA	r 8320 8320
	8321 8321	GAAATGTACTACGTGTCTGGAGCCCGCAGCAATGTCACATTTACTGTGAACCAAACATCCCGCCTCCTGATGAGGAGAA	8400 8400
	8401 8401	GAGGCGTCCAACTGGAAAAGTGACCCTGGAGGCTGACGTCATOCTCCCAATTGGGACACGCAGTGTTGAGACAGACAAGA	3 8480 8480

WN	02 x	M66 Variant ⇒ DNA Alignment	:
•	-	GACCCCTGGACAAGAGGCCCATAGAAGAAAGGGTTGAGAGGATAAAATCTGAGTACATGACCTCTTGGTTTTATGACAAT 8	3560 ···
	8481	GACAACCCCTACAGGACCTGGCACTACTGTGGCTCCTATGTCACAAAAACCTCCGGAAGTGCGGGGAGCATGGTAAATGG	3640
	8561	***************************************	,,,,
	8641	TGTTATTAAAATTCTGACATATOCATGGGACAGGATAGAGGAGGGTCACAAGAATGGCAATGACACACAC	0120,
	8721	GACAGCAAAGACTGTTTAAAGAAAAAGTTGACACCAGAGCAAAGGATCCACCAGCGGGAACTAGGAAGATCATGAAAGTT	0000
	8801	GTCAACAGGTGGCTGTTCCGCCACCTGGCCAGAGAAAAGAACCCCAGACTGTGCACAAAGGAAGAATTTATTGCAAAAGT	0000
	8881	CCGAAGTCATGCAGCCATTGGAGCTTACCTGGAAGAACAAGAACAGTGGAAGACTGCCAATGAGGCTGTCCAAGACCCAA	0300
	8961	AGTTCTGGGAACTGGTGGATGAAGAAAGGAAGCTGCACCAACAAGGCAGGTGTCGGACTTGTGTGTACAACATGATGGGG	304Ģ
	9041	ANAGAGAAGAAGCTGTCAGAGTTTGGGAAAGCAAAGGGAAGCCGTGCCATATGTATATGTGGCTGGGAGCCGGGTA	3120
	9121	TCTTGAGTTTGAGGCCCTGGGATTCCTGAATGAGGACCATTGGGCTTCCAGGGAAAACTCAGGAGGAGGAGGAAGGCA	3200
	9201	TTGGCTTACAATACCTAGGATATGTGATCAGAGACCTGGCTGCAATGGATGG	3260
	9281	GGATGGGACACGCGCATCACAGAGGCAGACCTTGATGATGAACAGGAGATCTTGAACTACATGAGCCCACATCACAAAAA	7300
	9361	ACTGGCACAAGCAGTGATGGAAATGACATACAAGAACAAAGTGGTGAAAGTGTTGAGACCAGCCCCCAGGAGGGAAAGCCT	7440
	9441	ACATGGATGTCATAAGTCGACGACACCAGAGAGGATCCGGGCAGGTAGTGACTTATGCTCTGAACACCATCACCAACTTG	9320
	9521	AAAGTCCAATTGATCAGAATGGCAGAAGCAGAGATGGTGATACATCACCAACATGTTCAAGATTGTGATGAATCAGTTCT	7600
	9601	GACCAGGCTGGAGCCATGGCTCACTGAGCACGGATGTGACAGACTGAAGAGGGATGGCGGTGAGTGGAGACGACTGTGTGG	9680 9680
		TCCGGCCCATCGATGACAGGTTCGGCCTGGCCCTGTCCCATCTCAACGCCATGTCCAAGGTTAGAAAGGACATATCTGAA	9760
		TGGCAGCCATCAAAAGGGTGGAATGATTGGGAGAATGTGCCCTTCTGTTCCCACCACTTCCATGAACTACAGCTGAAGGA	9840
	9841 9841	TGGCAGGAGGATTGTGGTGCCTTGCCGAGAACAGGACGAGCTCATTGGGAGAGGAAGGGTGTCTCCAGGAAACGGCTGGA	9920 9920
	9921	TGATCAAGGAAACAGCTTGCCTCAGCAAAGCCTATGCCAACATGTGGTCACTGATGTATTTTCACAAAAGGGACATGAGG	10000
	10001	CTACTGTCATTGGCTGTTTCCTCAGCTGTTCCCACCTCATGGGTTCCACAAGGACGCCACAACATGGTCGATTCATGGGAA	10080
•	1008	AGGGGAGTGGATGACCACGGAAGACATGCTTGAGGTGTGGAACAGAGTATGGATAACCAACAACAACCACATGCAGGACA	. 10160
	1016	AGACAATGGTGAAAAATGGAGAGATGTCCCTTATCTAACCAAGAGACAAGACAAGCTGTGCGGGATCACTGATTGGAATG	3 10240 10240

#### ### DNA Strider 13f7 ### Thursday, October 21, 2004 3:10:16 PM

WN02 M	Prot. x M66 M	Prot. => Prote	ein Alignment	•
Protein	sequence	.75 aa	SLIVOTEGESTL VVILILIVAPAYS	
Protein	sequence	75 aa	SLTVQTHGRSTL VVPILLVAPAYS	
Misma Gap r		Standard Single Bloc 6-aa Smaller (1) Medium (2) HLOSUM62		:
• : -	SLIVOTHGESTI	ANKKGAWMDSTK	40 60 KATRYLVKTESWILKNPGYALVAAVIGWALGSNIMORVVFVVIJLLVAPAYS KATRYLVKTESWILKNPGYALVAAVIGWALGSNIMORVVFVV LLLVAPAYS KATRYLVKTESWILKNPGYALVAAVIGWALGSNIMORVVFVVPIJLVAPAYS 40 60	75 75
8 Id	entity = 98	.7 (74/75)		

#### **CLAIMS**

- 1. A recombinant Flavivirus comprising a membrane protein mutation.
- 2. The Flavivirus of claim 1, wherein the mutation attenuates the Flavivirus.
- 3. The Flavivirus of claim 2, wherein the mutation decreases the viscerotropism/viremia of the Flavivirus.
- 4. The Flavivirus of claim 1, wherein the mutation results in increased stability of the Flavivirus, relative to a corresponding Flavivirus lacking the mutation.
- 5. The Flavivirus of claim 1, wherein the mutation results in increased virus replication in cells, relative to a corresponding Flavivirus lacking the mutation.
  - 6. The Flavivirus of claim 1, wherein the Flavivirus is a chimeric Flavivirus.
- 7. The Flavivirus of claim 6, wherein the chimeric Flavivirus comprises the capsid and non-structural proteins of a first Flavivirus and the membrane and/or envelope proteins of a second Flavivirus.
- 8. The Flavivirus of claim 7, wherein the first Flavivirus is a yellow fever virus.
  - 9. The Flavivirus of claim 8, wherein the yellow fever virus is YF-17D.
- 10. The Flavivirus of claim 7, wherein the second Flavivirus is a Japanese encephalitis virus.
- 11. The Flavivirus of claim 7, wherein the second Flavivirus is a West Nile virus.

12. The Flavivirus of claim 7, wherein the second Flavivirus is selected from the group consisting of a dengue virus, St. Louis encephalitis virus, Murray Valley encephalitis virus, and Tick-borne encephalitis virus.

- 13. The Flavivirus of claim 12, wherein the dengue virus is dengue-1, dengue-2, dengue-3, or dengue-4 virus.
- 14. The Flavivirus of claim 1, wherein the mutation is within the transmembrane domain of the membrane protein.
- 15. The Flavivirus of claim 14, wherein the mutation is a substitution in one or more amino acids corresponding to the region of amino acids 40-75 of the membrane helix within the membrane protein of a Japanese encephalitis virus or a West Nile virus.
- 16. The Flavivirus of claim 15, wherein the mutation is a substitution of an amino acid corresponding to amino acid 60 of the membrane protein of a Japanese encephalitis virus.
- 17. The Flavivirus of claim 16, wherein the mutation results in a substitution of arginine with cysteine at amino acid position 60 of the membrane protein.
- 18. The Flavivirus of claim 15, wherein the mutation is a substitution of an amino acid corresponding to amino acid position 66 of the membrane protein of a West Nile virus.
- 19. The Flavivirus of claim 18, wherein the mutation results in a substitution of leucine with proline at amino acid position 66 of the membrane protein.
- 20. The Flavivirus of claim 15, wherein the mutation is in one or more amino acids corresponding to those at positions 60, 61, 62, 63, 64, 65, or 66 of the membrane protein of Japanese encephalitis virus or West Nile virus.

21. The Flavivirus of claim 1, wherein the mutation is in the ectodomain of the membrane protein.

- 22. The Flavivirus of claim 21, wherein the mutation is in an amino acid selected from the group consisting of amino acids 1-5 of the ectodomain.
- 23. The Flavivirus of claim 22, wherein the mutation is a substitution in amino acid 5 of the ectodomain.
- 24. The Flavivirus of claim 23, wherein the mutation is a substitution of glutamine with proline.
- 25. The Flavivirus of claim 1, wherein the Flavivirus comprises one or more envelope protein mutations in residues corresponding to West Nile virus envelope protein amino acids selected from the group consisting of amino acids 107, 138, 176, 177, 224, 264, 280, 316, and 440.
- 26. The Flavivirus of claim 25, wherein the Flavivirus comprises envelope protein mutations in residues corresponding to West Nile virus envelope protein amino acids 107, 316, and 440.
- 27. The Flavivirus of claim 25, wherein the Flavivirus comprises mutations at residues corresponding to West Nile virus position 66 of the membrane protein and positions 107, 316, and 440 of the envelope protein.
- 28. The Flavivirus of claim 1, further comprising a mutation in the hydrophobic pocket of the hinge region of the envelope protein of the Flavivirus.
- 29. The Flavivirus of claim 28, wherein the mutation is present in an amino acid corresponding to amino acid 204 of the dengue 1 virus envelope protein.

30. The Flavivirus of claim 28, wherein the mutation is in one or more hinge region amino acids corresponding to yellow fever virus envelope protein amino acids 48-61, 127-131, and 196-283.

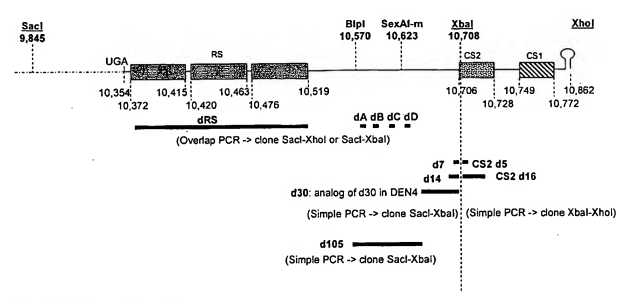
- 31. The Flavivirus of claim 1, further comprising an attenuating mutation in the 3'-untranslated region of the Flavivirus.
- 32. The Flavivirus of claim 1, further comprising an attenuating mutation in the capsid protein of the Flavivirus.
- 33. A vaccine composition comprising the Flavivirus of claim 1 and a pharmaceutically acceptable carrier or diluent.
- 34. A method of inducing an immune response to a Flavivirus in a patient, the method comprising administering to the patient the vaccine composition of claim 33.
- 35. The method of claim 34, wherein the patient does not have, but is at risk of developing, infection by the Flavivirus.
  - 36. The method of claim 34, wherein the patient is infected by the Flavivirus.
- 37. A method of producing a vaccine comprising a recombinant Flavivirus, the method comprising introducing a mutation into the membrane protein of the Flavivirus.
- 38. The method of claim 37, wherein the mutation attenuates the Flavivirus, relative to a corresponding Flavivirus lacking the mutation.
- 39. The method of claim 37, wherein the mutation results in increased stability of the Flavivirus, relative to a corresponding Flavivirus lacking the mutation.

40. The method of claim 37, wherein the mutation results in increased replication of the Flavivirus, relative to a corresponding Flavivirus lacking the mutation.

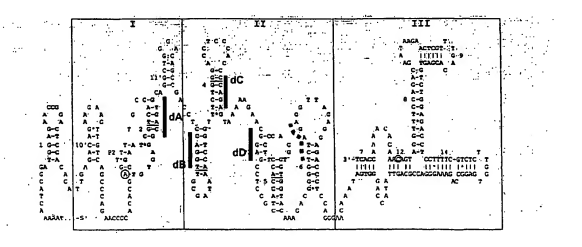
- 41. The method of claim 37, wherein the Flavivirus is a chimeric Flavivirus.
- 42. A nucleic acid molecule corresponding to the genome of the Flavivirus of claim 1 or the complement thereof.
- 43. A method of manufacturing the Flavivirus of claim 1, the method comprising introducing a nucleic acid molecule corresponding to the genome of the Flavivirus into cells and isolating Flavivirus produced in the cells from the cells or the supernatant thereof.
  - 44. The method of claim 43, wherein the cells are Vero cells.
- 45. The method of claim 43, wherein the cells are cultured in serum free medium.

Fig. 1. Deletions in the 3'UTR's of ChimeriVax<sup>TM</sup>-WN04-3'UTR candidates. Panel A: 3'UTR organization and introduced deletions. Panel B: predicted stem-loop structures (Proutski et al., J. Gen. Virol. 78:1543-1549, 1999) including ones that can be destabilized by small deletions dA – dD. Panels C and D, predictions of YF17D and dC mutant 3'UTR structures, respectively, by Zuker's algorithm.

A.



06132.099WO3 drawings.doc B.



Note: Dotted line in 2B designates XbaI restriction site at nucleotide 10,708.

Fig. 1. ...continued

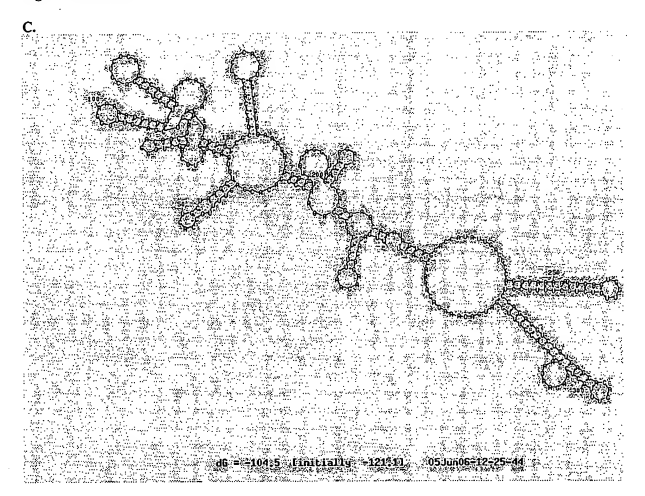


Fig. 1. ...continued

D.

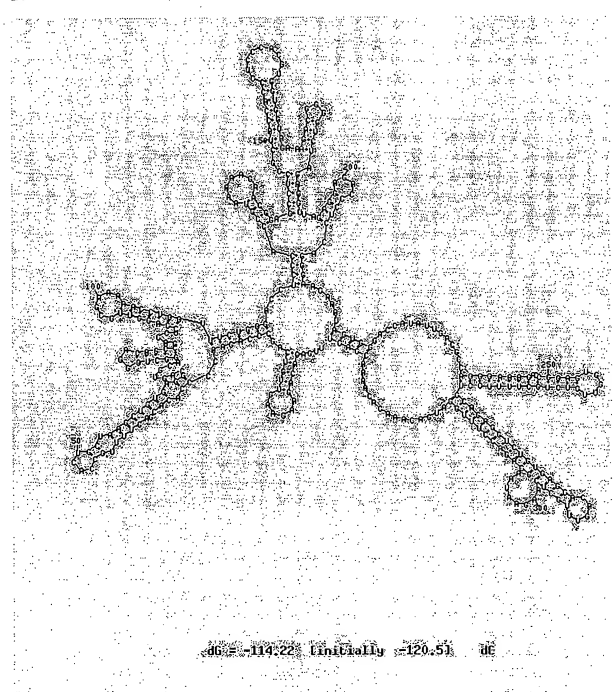
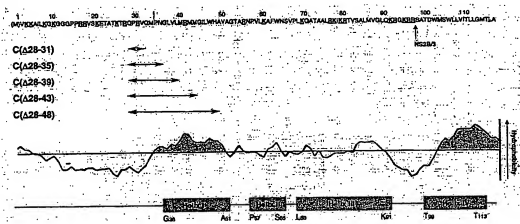


Fig. 2. Attenuating deletions for ChimeriVax<sup>TM</sup>-WN04-C variants (U.S. Patent Application Nos. 60/674,546 and 60/674,415). Panel A: previously described deletions introduced in the C protein of TBE virus (Kofler et al., J. Virol. 76:3534-3543, 2002). Panel B: computer-predicted structure of the YF 17D-specific protein C and proposed deletions C1-C5 (boxed) to be introduced in ChimeriVax<sup>TM</sup>-WN02.





## B.

C1 C2 C3 C4 C4 C5 C5 C5 C5 C6 C6 C7 C5 C6 C7 C5 C6 C7 C5 C7 C5 C7
Hydrophobic stretches
lace:
RSHDVLTVOFLILGELLHTGG  D sec HEHHHHHHHHHHHH  I sec G-Hellx.IV.
3 acc

Fig. 3. Replication in HepG2 cells.

## West Nile and YF/17D in HepG2 Cells MOI 0.005

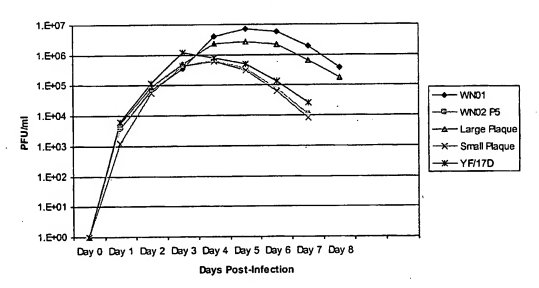


Fig. 4. Replication in THLE-3 cells.

## West Nile and YF/17D in THLE-3 Cells MOI 0.005

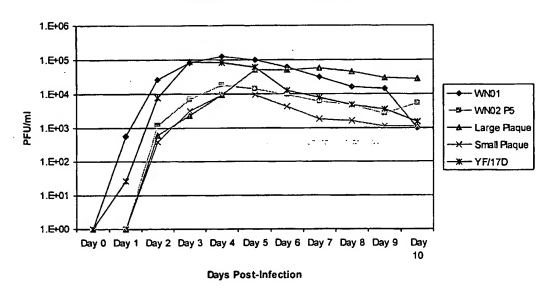


Fig. 5. Viremia in hamsters inoculated with ChimeriVax<sup>TM\_</sup>WN02 P5 (mixed plaque), S plaque (PMS, P10), or L plaque (PMS, P10) viruses.

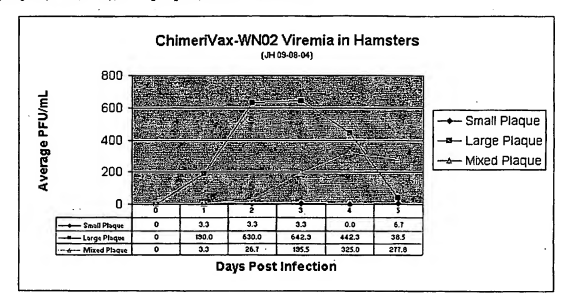


Fig. 6. Small-scale genetic stability passages (g.s.) from the uncloned P2 PMS virus in SF Vero cells.

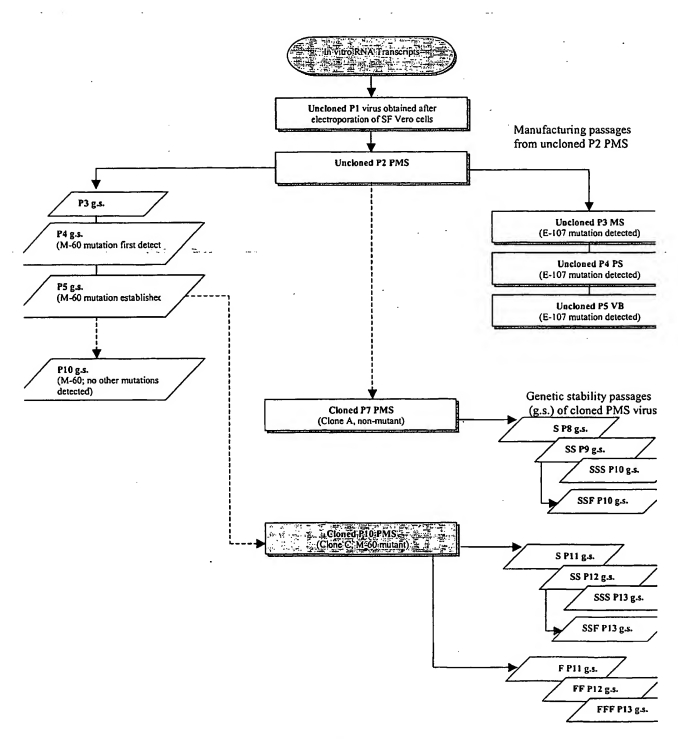


Fig. 7. A graph showing growth curves of SF ChimeriVax<sup>TM</sup>-JE viruses of the invention (uncloned P2, P3 MS (E-107), P4 PS (E-107), P5 g.s. (M-60), and P5 VB (E-107)) at the indicated times post-infection, which shows higher growth rates in SF culture of virus samples containing the M-60 and E-107 mutants as compared to non-mutant virus (P2).

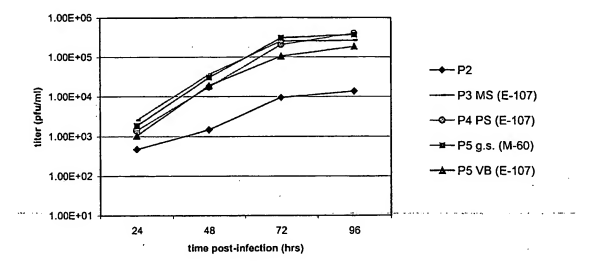


Fig. 8A. pH infectivity threshold analyses.

Fusion assay of
Clone E (M-5 mutant) compared to nonmutant Clone A P7,
Clone C P10 (M-60), and clone I (E-107) of Chimeri Vax-JE

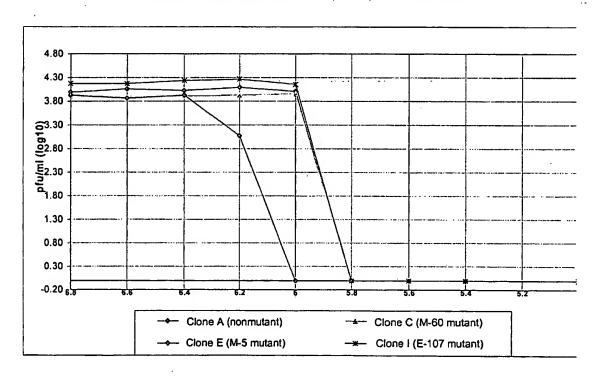


Fig. 8B. Survival Plot of ChimeriVax<sup>TM</sup>-JE vaccine (1.9 log<sub>10</sub> PFU/dose as determined by back titration of inocula) in comparison to ChimeriVax<sup>TM</sup>-JE M5 mutant (1.4 log<sub>10</sub> PFU/dose as determined by back titration of inocula) in 3-4 day old suckling mice inoculated by the intracerebral route.

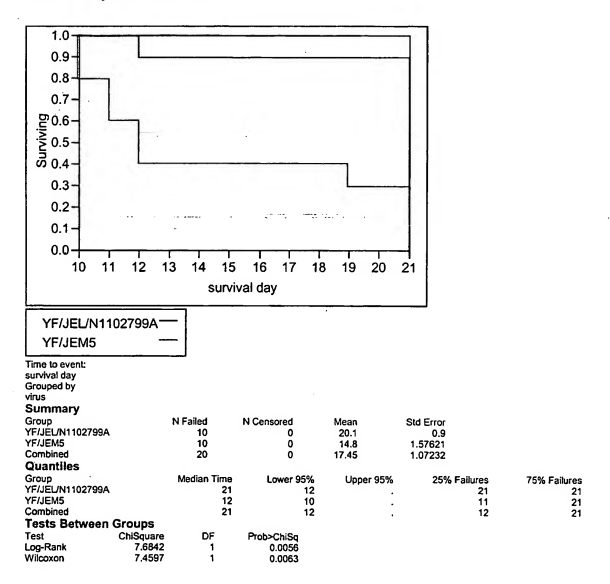


Fig. 8C. Survival Plot of ChimeriVax<sup>TM</sup>-JE M5 mutant virus (1.4 log<sub>10</sub> PFU/dose as determined by back titration of inocula) in comparison to YF-VAX® (0.9 log<sub>10</sub> PFU/dose as determined by back titration of inocula) in 3-4 day old suckling mice inoculated by the intracerebral route.

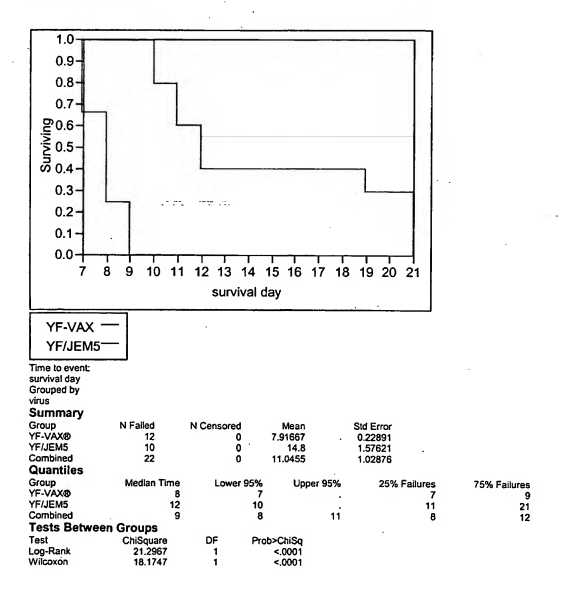


Fig. 8D. Indirect Fusion assay: comparison P7 and P10 of ChimeriVax<sup>TM</sup>-DEN1-4 viruses. The virus output for each experiment was determined by standard plaque assay. A, ChimeriVax<sup>TM</sup>-DEN1 PMS P7 (triangles) and P10 (diamonds); B, ChimeriVax<sup>TM</sup>-DEN2 PMS P7 (triangles) and P10 (diamonds); C, ChimeriVax<sup>TM</sup>-DEN3 PMS P7 (triangles) and P10 (diamonds); D, ChimeriVax<sup>TM</sup>-DEN4 PMS P7 (triangles) and P10 (diamonds)

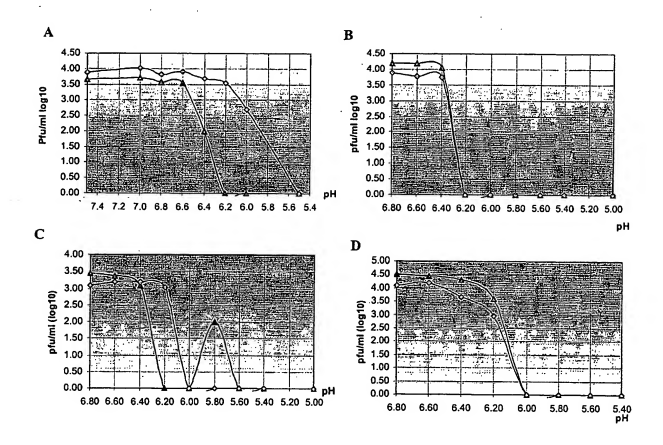


Fig. 8E. Indirect Fusion assay with the ChimeriVax<sup>TM</sup>-DEN3, comparing the PMS (P7) vaccine with the Vaccine lot (P10) and the P15 virus.

The virus output for each experiment was determined by standard plaque assay. ChimeriVax<sup>TM</sup>-DEN3 PMS P7 (triangles), P10 (diamonds), and P15 (squares).

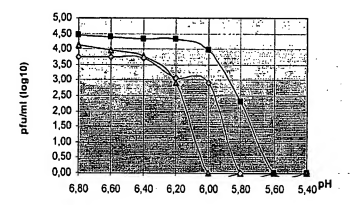


Fig. 8F. Structure of DEN1 E-protein dimmer (aa 1 to 394) of ChimeriVax<sup>TM</sup>-DEN1 virus (Guirakhoo et al., J.Virol. 78:9998-10008, 2004). (A) The position of the positively charged lysine (K) at residue 204 of the P7 (PMS, 204K) virus is shown by CPK (displays spheres sized to van der Waal radii) representation. Three structural domains are shown in medium grey (domain I), light grey (domain II), and dark grey (domain III). (B) Close up of marked area in panel A. (C) The same area as in panel B from the E protein model of the mutant DEN1 virus (P10, 204R shown in red). Selected amino acids in panel B and C are shown in stick representation. Grey, carbon; blue, nitrogen; red, oxygen; yellow, sulfur.

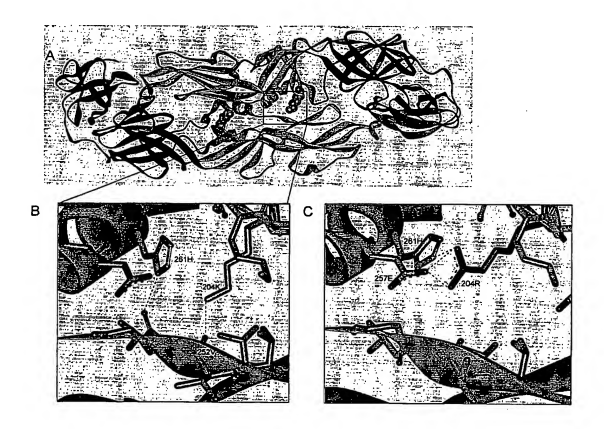


Fig. 9A. A graph showing the penetration efficiency of ChimeriVax<sup>TM</sup>-JE viruses M60 mutant (Clone C), E107 mutant (Clone I), and non-mutant (Clone A) at the indicated times. These results indicate that the M60 mutation facilitates penetration in SF Vero cells apparent at the 5 and 10 minute time points. SF Vero cells were infected with appropriately diluted viruses (Clones A, C, and I in serum free medium) for 5, 10, 20, or 60 minutes, and then were treated for 3 minutes with a solution of 0.1 M glycine, 0.1 M NaCl, pH 3.0, to inactivate extracellular virus. Wells were washed twice with PBS and then monolayers were overlaid with methyl-cellulose followed by staining plaques on day 5 with crystal violet. Efficiency of penetration is shown as percentages of observed plaque numbers after glycine treatment as compared to control infected wells that were treated with PBS instead of glycine.

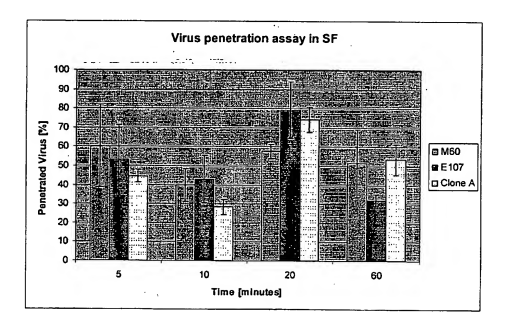


Fig. 9B. Schematic representation of the location of the E-107, M-5, and M-60 amino acid residues in the envelope proteins E and M, illustrating the hypothetical effect of the M-5 residue on fusion. The dashed stretch at the tip of domain II of the E protein containing the E-107 residue represents the fusion peptide (c-d loop), which inserts into the cell membrane (Rey et al., Nature 375:291-298, 1995). The M-5 residue is at the N terminal of the ectodomain part of the M protein. The E protein monomers rearrange into trimeric complexes, which fold to force the cell and virus membranes to fuse (Modis et al., Nature 427(6972):313-319, 2004). M may be a functional component of the complexes, e.g., facilitating fusion of the viral membrane with the cell membrane via its interaction with the E protein. The M-60 residue is between the two C-terminal transmembrane stretches of M and may participate in the interaction of the cell and viral membranes during fusion.

